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oligene GmbH, Berlin/DE

(vormals: Pathoarray GmbH)

Bezeichnung:

Nukleinsäurearray

IPC:

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Die angehefteten Stücke sind eine richtige und genaue Wiedergabe der ursprünglichen Unterlagen dieser Patentanmeldung.

> München, den 2. Juli 2003 **Deutsches Patent- und Markenamt** Der Präsident

Im Auftrag

rofsky

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Belegenemplar Barf nicht geändert werden

Beschreibung

Die Erfndung betrifft einen Array bestehend aus Oligooder Polynukleotidsonden, die immobilisiert auf einen festen Träger aufgebracht sind. Das Array ist dadurch charakterisiert, dass auf der Oberfläche Sequenzen einer Auswahl oder aller der in den Tabellen 1-6 genannten selektiven Monozyten-Makrophagen-Gene gebunden sind. Dieser Nukleinsäure-Array ermöglicht die Diagnose der rheumatoiden Arthritis, eine begleitende Analyse der Behandlungseffektivität und die Überwachung von Nebenwirkungen bei der anti-Tumornekrosefaktor-(TNF)-Therapie und somit die Auswahl der für den jeweiligen Patienten mit rheumatoider Arthritis am wirkungsvollsten Therapie. Die vorliegende Erfindung betrifft ferner einen Nukleinsäure-Array zur Prognose und zur Entwicklung neuer anti-TNF gerichteter Pharmaka oder solcher Pharmaka, die in dessen Regelkreis eingreifen.

Die Zellen des Monozyten / Makrophagen-Systems sind an der Aktivierung und Aufrechterhaltung von Entzündungskaskaden im Blut und im Gewebe z. B. im Rahmen der rheumatoiden Arthritis und bei anderen chronisch entzündlichen Erkrankungen, aber auch bei autoaggressiven Erkrankungen wesentlich beteiligt. Bei diesen Erkrankungen sind Monozyten und Makrophagen hoch aktiviert, zeigen Veränderungen im Besatz ihrer Oberflächen-Moleküle, treten mit anderen Zellen in Kontakt und sezernieren bestimmte Botenstoffe wie u. a. TNF-alpha, die dafür sorgen, den Entzündungsvorgang zu unterhalten. TNF-alpha ist ein von Monozyten / Makrophagen, Lymphozyten und Mastzellen gebildetes Zytokin mit Einfluss auf Entzündung, Sepsis, Lipidund Proteinstoffwechsel, Blutbildung, Angiogenese, Wund-

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heilung und Immunabwehr, das aber auch zytolytische bzw. zytostatische Wirkung auf Tumorzellen hat.

Bei entzündlichen Erkrankungen zeigen Monozyten

Makrophagen ein charakteristisches, pathologisch verändertes Genexpressionsmuster mit deutlichen Abweichungen im Vergleich zu gesunden Probanden. Mit dem Fachmann bekannten bioinformatischen Methoden wie z. B. der Signifikanz- und Clusteranalyse lassen sich u. a. Gene mit ähnlichem Verhalten und hoch- oder niederregulierte Gene aus den Hybridisierungsmustern eines Nukleinsäurearrays bestimmen.

Die zunehmende Verfügbarkeit der Hochdurchsatz-Verfahren in Form von Nukleinsäurearrays, die exponentiell anwachsenden Informationen zum humanen Genom und der Genexpression, sowie die globale Vernetzung von Datenbanken mit strukturierten biomedizinischen Informationen wird die Betrachtungsweise chronisch entzündlicher und entzündlich-rheumatischer Krankheitsbilder grundlegend verändern. Aus dem verbesserten Verständnis der molekularen Grundlagen der zell-, gewebs- und krankheitsspezifischen Genexpression lassen sich die molekularen Abläufe definieren und tragen dazu bei, eine frühere Diagnose und verbesserte Prognose zu erlauben. Zum anderen gewährleisten Mikroarray-Technologien effektivere Therapieformen für die rheumatoide Arthritis und für andere chronisch entzündliche Erkrankungen zu entwickeln und ermöglichen ein schnelles Screeningsystem. Ferner erlauben diese multiplen Verfahren die Entwicklung von pharmazeutischen und biologisch wirksamen Medikamenten (Biologicals) zu beschleunigen und die Testphasen der Medikamentenwirkung, wie auch die Beurteilung der Medikamenten Nebenwirkungen schneller beurteilen zu können. Aus diesem Grund stellt

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dieses Verfahren einen volkswirtschaftlichen und wirtschaftlichen Gewinn dar.

Die Mikroarray Technologie stellt eine Miniaturisierung analytischer Verfahren auf der Basis der DNA- bzw. RNA-Hybridisierung im Hochdurchsatz-Verfahren dar. Gleichzeitig können dadurch viele tausend verschiedene DNA/DNA-(DNA/RNA-) Wechselwirkungen innerhalb eines Testansatzes analysiert werden. mRNA-Expressionsprofile werden mittels DNA-Arrays durch die Hybridisierung von markierten cRNA oder cDNA-Proben bestimmt. Diese Technologien erfordern ein hohes Maß an Automatisierung und Standardisierung mit Aufbau und Nutzung entsprechender Proben- und Datenbanken (Sequenzinformationen, Oligonukleotide). Die derzeit verwendeten DNA-Arrays unterscheiden sich im Trägermaterial Glasoberflächen, Edelmetall bedampfte (Nylonmembranen, Glasoberflächen, Kunststoffe), der Länge bzw. der Herstellung der an den Träger immobilisierten DNA-Sequenzen und der Markierungstechnik für eine zu bindende Probe. In Analogie zu den Methoden der DNA-Hybridisierung beim Southern-/Dot-Blot können DNA-Sequenzen auf einem Filter punktförmig und in systematischer Reihenfolge mit einem Piezo-Druckverfahren durch Spotting, durch Druckkopf durch Photolithographie (Tintenstrahltechnologie) oder (chemische Direktsynthese auf dem Trägermaterial) fixiert werden. Die DNA kann dabei eine cDNA, ein PCR-Produkt oder ein synthetisch hergestelltes Oligonukleotid sein. Jede dieser aufgetragenen Sequenzen ist damit einem spezifischen Ort in einer bekannten Anordnung zugeteilt. Aus einer klinischen oder aber pharmazeutisch zu untersuchenden Probe kann RNA aufgereinigt werden und nach Umschreibung durch reverse Transkription mit den auf dem Array befindlichen komplementären Nukleinsäurensträngen die in einer hohen genomweiten Anzahl oder aber einer bereits

Streptavidin Wechselwirkungen, Digoxigenin-Enzym Verstärkungen oder aber über direkte oder indirekte eingebaute Fluoreszenzfarbstoffe. Das Auslesen der Information erfolgt über die Intensität der Radioaktivität oder der Fluoreszenz an einem spezifischen Ort des Trägermaterials und lässt somit Rückschlüsse zu, welche relative Menge an spezifisch gebundener DNA- bzw. RNA-Sequenz in der markierten Probe vorhanden war.

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Das An- und Abschalten von Genen ist Grundlage aller biologischen Prozesse und außerdem eine extrem sensitive Antwort auf veränderte äußere Bedingungen. Mit der Extraktion von RNA aus einer biologischen Probe, dem Einwirken von markierter cDNA oder RNA auf einen Nukleinsäure-Array (Hybridisierung) und dessen Analyse ist innerhalb kürzester Zeit eine große Fülle von Informationen über den Zustand der Zellen in der biologischen Probe unter veränderten Bedingungen möglich. Die auf der Hybridisierung von Nukleinsäuren beruhende Technologie hat den Vorteil einer extrem hohen Spezifität, Sensitivität und relativ leichten, schnellen Durchführbarkeit.

Geschieht das An- oder Abschalten von Genen in Monozyten/Makrophagen in nicht physiologischer Weise, so kann es die Ursache von entzündlichen Erkrankungen oder ein messbares Zeichen für diese sein. Die Therapie mit anti-TNF wirksamen Medikamenten sollte im Idealfall die pathologisch veränderte Genexpression in den betroffenen Zellen auf das Niveau von gesunden Patienten normalisieren.

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Durch Untersuchung der Genexpressionsprofile ist zu erwarten, dass eine neue molekulare Charakterisierung der rheumatoiden Arthritis und anderer chronisch entzündlicher Erkrankungen möglich wird und damit eine Einteilung in Subgruppen nach pathophysiologischen Besonderheiten erfolgt. Bei den entzündungshemmenden anti-TNF Therapien stehen somit prognostische Vorhersagen in Aussicht über die Agressivität im weiteren Verlauf. Dies würde bereits frühzeitig Einfluß auf die Wahl und Intensität der medikamentösen Therapie mit den bisher bekannten bei chronischen Entzündungen verwendeten Medikamenten, aber auch mit biologisch wirksamen TNF-Blockern ausüben. Zum anderen ergeben sich hieraus weitere Ansatzpunkte, um die Therapieform im Hinblick auf die potentiellen Nebenwirkungen durch Einflussnahme dieser Medikamente zu gestalten und die Auswirkung der Nebenwirkungen rechtzeitig abzuschätzen.

Durch anti-TNF gerichtete Therapien bei der rheumatoiden Arthritis und anderen chronisch entzündlichen oder autoaggressiven Erkrankungen wird zum einen eine potentielle Entstehung neoplastischer Veränderungen bis hin zur Tumorbildung diskutiert, zum anderen vermindert die anti-TNF Therapie die Immunabwehr, sodass bei den behandelten Patienten vermehrt Infektionen auftreten, u. a. Tuberkulose.

Mit Hilfe von Nukleinsäure-Array-Systemen kann die Expression tumorrelevanter Gene im Verlauf der anti-TNF Behandlung überprüft und somit frühzeitig Hinweise auf mögliche neoplastische Veränderungen geben, so dass einer beginnenden Tumorentwicklung rechtzeitig entgegengesteuert und die anti-TNF Therapie entsprechend angepasst oder falls nötig abgebrochen werden kann.

Der Erfindung liegt die Aufgabe zugrunde, Mittel zur Überwachung der Wirksamkeit sowie von Nebenwirkungen der anti-TNF Therapie zu schaffen, aber auch die Feindiagnos-

tik einer entzündlichen Erkrankung und damit die Auswahl der für den jeweiligen Patienten effektivsten Therapieform zu ermöglichen. Eine weitere Aufgabe der vorliegendie Wirksamkeit und Nebenden Erfindung besteht darin, wirkungen neuer anti-TNF gerichteter Pharmaka im Rahmen von klinischen Studien zu verfolgen. Erfindungsgemäß wird ein neuer Array geschaffen bestehend aus Oligo- oder Polynukleotidsonden, die immobilisiert auf einem festen Träger aufgebracht sind. Verglichen mit bisher bekannten genomweiten DNA-Chips ist der Vorteil der Erfindung eine Kostenersparnis bei der Herstellung des Nukleinsäurearrays, weil es überwiegend nur Gene enthält, die zur Lösung der Aufgabe der Erfindung interessant sind, was den Aufwand der Datenauswertung minimiert und damit verbilligt.

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Erfindungsgemäß wird die Aufgabe durch einen Nukleinsäure-Array gelöst, auf dessen Oberfläche Sequenzen einer
Auswahl oder aller der in den Tabellen 1 bis 6 genannten
selektiven Monozyten-Makrophagen-Gene aufgebracht sind.
Anhand des Gen- oder Sequenznamens oder der AccessionNummer kann die Sequenz aus öffentlich zugänglichen Datenbanken, vorzugsweise GeneBank oder EMBL, ermittelt
werden. Die Sequenzen der aus dem Array befindlichen Nukleinsäuren können aus Genen bestehen, deren Expressionsniveau durch eine anti-TNF wirksame Therapie verändert
wird.

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Gegebenenfalls können auf dem erfindungsgemäßen Nukleinsäure-Array weitere Gene vorhanden sein, vorteilhaft solche, von denen bekannt ist, dass sie in jeder Zelle exprimiert werden und zur Grundausstattung der Zelle gehören. Die Gene, die für diese Nukleinsäuren codieren, werden üblicherweise als Haushalts- oder Housekeeping-

Gene bezeichnet und werden zur Normierung der erhaltenen Signale verwendet. Das Array kann die genannten Sequenzen in Form von DNA, komplementärer RNA oder chemisch modifizierten Nukleinsäuren, vorzugsweise PNA (protein nucleic acid) enthalten.

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Bei den Genen oder Gensequenzen kann es sich um krankheits- und nebenwirkungsrelevante selektionierte Gene der
rheumatoiden Arthritis oder anderer chronisch entzündlicher Erkrankungen handeln, vorzugsweise aus dem Monozyten/Makrophagen-Zellsystem. Gegebenenfalls können auf
den Oberfläche des Arrays auch Allele, Derivate und/oder
Splicingvarianten der Gen- oder Genteilsequenzen oder Oligomersequenzen vorliegen. Die Übereinstimmung der Sequenzen auf dem Array mit den entsprechenden Sequenzen in
Tabelle 1-6 soll dabei mindestens 80 % in den Proteinkodierenden Abschnitten der mRNA betragen.

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Der Träger, auf den die Nukleinsäuren aufgetragen werden, kann jeder Träger sein, der normalerweise für RNA- oder DNA Arrays verwendet wird. Die Verfahren zum Auftragen und Immobilisieren der Nukleinsäuren sind Stand der Technik und dem Fachmann bekannt. Zur Kopplung der genannten Sequenzen kann der Träger mit reaktiven Gruppen, Metallverbindungen oder Legierungen beschichtet sein. Die Gene oder Gensequenzen können bespielsweise durch Spottingverfahren, Immobilisierungsverfahren oder durch in-sito Syntheseverfahren von Oligomeren oder spiegelbildlich in Form von RNA aufgebracht werden.

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Das erfindungsgemäße Array kann beispielsweise zur Messung der Monozyten/Makrophagen Aktivierung oder der Entzündungsaktivität im Blut oder Zellgewebe bei entzündlichen Erkrankungen, vorzugsweise der rheumatoiden Arthri-

tis verwendet werden. Das Array kann z. B. zur Früherkennung der genannten Erkrankungen bei genetisch vorbelasteten Patienten verwendet werden, noch bevor sich klinische Symptome manifestieren. Ein weiterer Einsatzbereich ist die Feindiagnostik, vorzugsweise die Einteilung von Patienten in Subgruppen, die jeweils eine unterschiedliche Therapie und unterschiedliche Medikamente benötigen. Das Array kann ferner zur Therapieüberwachung, zur Verfolgung von Nebenwirkungen, zur Erstellung einer Prognose und zur Indentifizierung neuer pharmazeutischer Targets bei den genannten Erkrankungen verwendet werden.

Dazu werden den zu untersuchenden Patienten Blut oder Gewebeproben entnommen, aus denen RNA mit bekannten Standardtechniken isoliert und gegebenenfalls als Gesamt-RNA oder Poly A+-RNA weiterverwendet wird. Mit reverser Transkriptase kann die RNA in cDNA umgeschrieben und dabei mit einer Markierung versehen werden, z. b. einem Fluoreszenzfarbstoff, einem radioaktiven Nuklid oder einem Enzym wie alkalische Phosphatase. Daneben kann die RNA direkt markiert oder unmarkiert zur Hybridisierung des Nukleinsäure-Arrays eingesetzt werden. Nach Hybridisierung des Arrays mit den Nukleinsäureproben und nachfolgenden Waschschritten kann die Bindung der Probe an die auf dem Array befindlichen Sequenzen mit jedem geeigneten Verfahren analysiert werden. Im Falle einer Fluoreszenzmarkierung sind dies optische Verfahren, bei radioaktiv markieren Proben käme eine Autoradiographie zur Anwendung und bei einer Enzymmarkierung entzymatische

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Nachweisverfahren, z.B. die Umsetzung eines farblosen Substrates zu einem farbigen Produkt.

Ein inverser Nachweis von festphasengebundener Total- oder mRNA mit den Sequenzen aus Tabelle 1-6 ist ebenfalls möglich. Dazu werden auf den RNA-Mikroarrays Blut- oder gewebsspezifische RNA-Moleküle von bis zu 500 Patienten gebunden. Der qualitative / quantitative Nachweis der Transkriptmenge relevanter Gene erfolgt dann mit den in Tabelle 1-6 beschriebenen selektionierten Genen, Genabschnitten oder Oligomeren. Die RNA-Proben werden auf Kopplungsträger gespottet und setzen sich aus Total-RNA oder messenger-RNA zusammen. Die RNA dient dabei als Target für die aus DNA-Mikroarrays abgeleiteten hoch signifikant exprimierten Gene nach Tabelle 1-6, die als markierte Sonden zur Hybridisierung eingesetzt werden. Vorgeschlagen wird das Koppeln biotinylierter RNA oder messenger-RNA auf Streptavidin beschichteten Glasträgern (Slides). Nach Markierung der RNA mit Biotinderivaten, wird die RNA auf Poly-L-Lysin behandelten vorzugsweise aber auf mit Streptavidin beschichteten Glas- oder Plastikslides durch Spotting aufgebracht und getrocknet. Eine Degradation der RNA wird so verhindert. Alternativ bietet sich eine kovalente Kopplung der RNA durch Bindung an reaktive Trägermaterialien an, die vorzugsweise durch UV-Bestrahlung katalysiert wird. Zusätzlich ist eine multiple, gleichzeitige Markierung verschiedener Gene, Geneinheiten oder Oligomere mit verschiedenen Markierungs-Spezies, z.B. Radioaktivität, Fluoreszein, Digoxigenin und enzymatischen Markierungen vorteilhaft.

Parallel unterschiedliche Markierungen der Sonden mit unterschiedlichen Fluoreszenzfarbstoffen sind möglich. Alternativ sind enzymatische oder aber radioaktive Sonden-

markierungen zu nennen. Zur Quantifizierung und Qualitätskontrolle werden markierte Haushaltsgene (alpha-, beta, gamma-Aktin, GAPDH usw.) eingesetzt. Bevorzugt wird der Nachweis hier parallel und gleichzeitig mit maximal 50

5 Gensonden pro Ansatz gleichzeitig durchgeführt.

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Neben der Vereinfachung der biometrischen Analyse durch Kopplung von RNA Spezies an Trägermaterialien erlaubt dieses System eine schnelle Diagnostik und bietet eine komplexe für den Patienten individuell schnelle Diagnostik, Prognostik und Therapiesteuerung. Insbesondere bei pharmakologischen Entwicklungsstrategien erlaubt das System eine schnelle Durchführung mit hohem Durchsatz.

Die folgenden Beispiele und Abbildungen dienen nur zur Erläuterung und beschränken in keiner Weise den Umfang der Erfindung.

1. Isolierung von Monozyten

Im hier angewandten Verfahren wurde die Auswahl selektiver hochreiner Monozyten des peripheren Blutes benutzt, um eine Aussage 1.) zur Krankheitsspezifität, 2.) der Anwendung des Therapeutikums anti-TNF-alpha, als "Biological", 3.) im Vergleich zum Gesunden Probanden, als auch 4.) zur Bewertung von anti-TNF-alpha relevanten gendiagnostischen Möglichkeiten, zu ermöglichen. Dabei wurden die peripheren Blut-Leukozyten aus peripherem Blut durch eine Fikollgradienten-Dichtezentrifugation angereichert. Diese Fraktion, die individuell unterschiedliche Zusammensetzung aus Monozyten (5-12%), CD4+ T-Zellen (85-92%), CD8+ T-Zellen (5-10%), NK-Zellen (2-5%), basophilen und neutrophilen Granulozyten aufweist, wurde zur Gewinnung spezifischer Monozytenfraktionen weiteren Reinigungsschritten unterzogen. Hierbei kamen sowohl Negativ-

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selektionen, bei denen sämtliche andere Zellfraktionen über magnetische Beads-Antikörper Wechselwirkungen entfernt werden, als auch Positivselektionen durch CD14+Markierung über magnetische Beads oder aber FACS Zellsortierungsverfahren zum Einsatz. Bei beiden Verfahren ergaben sich Monozyten-Zellreinheiten von ca. 96 %.

2. RNA-Gewinnung

Die reinen Monozytenfraktionen wurden in RNA-Lysepuffer aufgenommen und die RNA dann über einen kommerziell erhältlichen RNA Reinigungskit (Qiagen) gereinigt. Die RNA wurde über etablierte cDNA Umschreibemethoden durch reverse Transkription in cDNA umgeschrieben und dann einem weiteren linearen Amplifikationsschritt durch das angewandte "Eberwine Protokoll" zur Herstellung von aRNA (amplifizierte RNA) unterzogen. Die Quantität und Qualität der RNA, cDNA, und aRNA wurde jeweils durch Gelelektrophorese, photometrische Bestimmung und über Messungen mit dem Bioanalyzer 2100 (Fa. Agilent) verifiziert.

3. Affymetrix Chip Hybridisierung

Für Expressionsanalysen werden im System der Firma Affymetrix spezifische direkt aus Datenbanksequenzen abgeleitete Oligonukleotide als DNA-Proben verwendet. Diese werden auf dem Array mit Targets aus fluoreszenz-markierten revers transkribierten Proben in Form von cDNA oder mit linear amplifizierten Proben in Form von aRNA hybridisiert.

Die Hybridisierung des genomweiten Affymetrix-Arrays (U133A) und weitere Bearbeitung erfolgt maschinell unter
Standardbedingungen nach Angaben des Herstellers Affymetrix in einem speziellen Hybridisierungs- und Waschgerätgerät mit den speziellen Puffern. Genexpressionsmuster

werden nach Hybridisierung über das Verhältnis der Fluoreszenzintensitäten bei einer bestimmten Wellenlänge erstellt. Solche Hochdurchsatz-Expressionsanalysen erlauben
Vergleiche der Expressionsmengen von Genen gleichzeitig
in gesundem und krankem Personen oder Vergleiche der Ge-

in gesundem und krankem Personen oder Vergleiche der Genexpression vor und nach Arzneimittelzugabe zur Risikoabschätzung (Pharma-/Toxikogenomik), zur Feindiagnostik und Abschätzung der Komplexität von Erkrankungen.

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4. Datenauswertung

Zum Einsatz kamen dabei aRNA Proben aus peripheren BlutMonozyten 1.) gesunder Blutspender, 2.) chronisch aktiver
Patienten mit rheumatoider Arthritis vor Behandlung und
3.) nach Behandlung mit TNF-alpha Antikörpern. Der Behandlungserfolg wurde über laborklinisch eindeutige Parameter und nach den klinisch anzuwendenden Kriterien der
internationalen gültigen Parameteruntersuchungen (ACRKriterien) abgeschätzt. Ziel und Zweck dieser Dreigruppenuntersuchung war es, charakteristische Genexpressionen
in folgenden Gruppendefinitionen festzustellen:

- 1.) Eine genregulatorische Krankheitsspezifität bei der aktiven unbehandelten rheumatoiden Arthritis, im Vergleich zur Genexpression gesunder Probanden.
- 2.) Eine genregulatorisch spezifische Interpretation der anti-TNF-alpha-Behandlung zu charakterisieren und eine Bewertung der Behandlung im Vergleich zur Genexpression der aktiven unbehandelten Krankheit und im Vergleich zur Genexpression der gesunden Probanden durchzuführen.

3.) Die Bewertung von Nebenwirkungen durch das Medikament="Biological" anti-TNF zu gewährleisten.

Hierbei wurde die spezifische Genexpression der anti-TNF-alpha behandelten Patienten mit rheumatoider

Arthritis mit der Genexpression der unbehandelten selben Patienten, und der von gesunden Blutspender verglichen.

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- Die Bearbeitung und Messung der einzelnen Genexpressionen innerhalb des genomweiten humanen Affymetrix-Arrays (U-133A) erfolgte innerhalb des zugehörigen Affymetrix Hybridisierungs-/ Wasch- und Auslesegerät System. Die Auswertung vollzieht sich in 4 Schritten:
 - 1. Bestimmung der bei der Expressionsanalyse detektierten signifikanten Gene, z.B. durch die "Fold-Change Method" oder SAM ("Significance Analysis of Microarrays").
- Separation der signifikanten Gene in verschiedene Sub-Populationen auf der Grundlage der Untersuchung der Expressionseigenschaften dieser Gene mittels Cluster-Analyse mit Verfahren wie "Hierarchical Clustering", "Self-Organizing Maps" oder "k-Means-Clustering".
 - 3. Auswertung des Verhaltens der signifikanten Gene innerhalb der Cluster unter Einbeziehung der klinischen Informationen (rheumatoide Arthritis (RA), anti-TNF-Therapie) und nach den Erfahrungswerten von
 Spezialisten.

4. Zuordnung der beteiligten Gene nach biologischen Pathways.

Allgemeines Verhalten der signifikanten Gene innerhalb der Cluster:

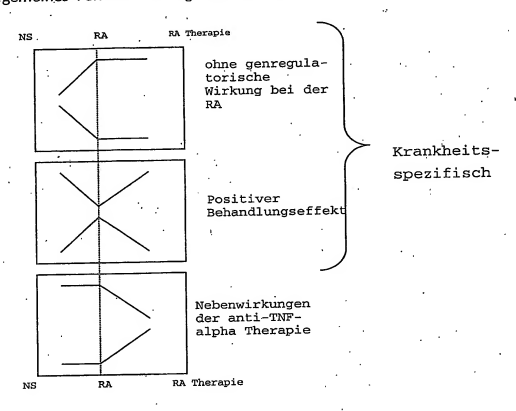


Abb. 1: schematische Darstellung der Clusteranalyse

Das Genexpressionsverhalten eines gesunden Normalspenders (NS) sowie und eines aktiven Patienten mit rheumatoider Arthritis (RA) vor und nach einer anti-TNF-alpha Therapie wurden mittels Clusteranalyse verglichen. Die Ergebnisse sind in den Abbildungen 1 und 2 dargestellt.

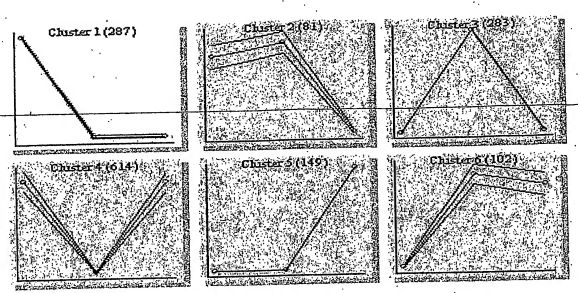


Abb. 2: Clusteranalyse anhand realer Daten.

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Dargestellt sind die Genexpressionen der Clusteranalyse (n=6 Cluster). Die Anzahl der beteiligten Gene ist in Klammern wiedergegeben. Als Ergebnis der Clusteranalyse erhält man zusätzlich zum durchschnittlichen Genexpressions-Verhalten aller in einem Cluster befindlichen Gene ein Vertrauensintervall.

Die Cluster weisen dabei folgende Charakteristiken auf:

CLUSTER-1: Die krankheitsspezische Genexpression ist kleiner im Vergleich zum Gesunden, die antiTNF-Behandlung ist hier ohne genregulatorische Wirkung.

CLUSTER-2: Nebenwirkungen: Dargestellt durch die Medikamentenwirkung der Anti-TNF-alpha Behandlung besteht eine verminderte Expression der zugehörigen Gene beim behandelten Patienten.

CLUSTER-3: Die krankeitsspezifische Genexpression größer im Vergleich zum Gesunden. Die anti-TNF-alpha Behandlung zeigt einen positiven Effekt.

CLUSTER-4: Die krankheitsspezifische Genexpression ist kleiner im Vergleich zum Gesunden. Die anti-TNF-Behandlung zeigt einen positiven Effekt.

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CLUSTER-5: Nebenwirkungen: Dargestellt dürch die Medikamentenwirkung der anti-TNF-alpha Behandlung besteht eine erhöhte Expression der zugehörigen Gene beim behandelten Patienten.

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CLUSTER-6: Die krankheitsspezische Genexpression ist größer im Vergleich zum Gesunden. Die anti-TNF-alpha Behandlung ist hier ohne genregulatorische Wirkung.

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In den Tabellen 1-6 sind die in den oben beschrieben Clustern enthaltenen Gene zusammen mit der Affymetrix Bezeichnung (links) und ihrer definierten GeneBank-Accession Nummer inkl. einer Beschreibung aufgeführt.

Tabelle 1: Gene aus Clusteranalyse 1

	1
Affymetrix-	i D J
interne Be-	
zeichnung	
211654 v at	gb:M17565.1 / DEF=Human MHC class 11 DQ-Deta associated with Zano, Zano 22 DB XREF=qi:188188 /FL=gb:M17565.1
200112	Consensus includes gb:X00452.1 /DEF=Human mRNA for DC classII histocompatibility antigen alpha-chaun. /FbA-mANA / PBA-B-ClassII histocompatibility antigenalpha-chain /DB_XREF=gi:32265 /UG=HS.198253 major histocompatibility complex,
213831_at	class II. Do alpha 1
100000	b:BF338947 /FEA=EST /DB_XREF=g1:1128530/ /DP_ron induced transmembrane protein 3 (1-80)
212203_X_ac	oc-ne.1022 gb:MIG272 gb:MIG272 / IPEE-Human MHC class II HIA-DR2-DW12 mRNA DGW1-beta, complete cds. /FEA=mRNA /GEN=HIA-DRB2 / DB XREFeq:188397 /UG-Hs.73931 major histocompatibility complex, class II, DQ beta 1 /FL=gb:M60028.1 gb:M17564.1
209480_at	gb:H81140.1 gb:M81141.1 gb:M16276.1 gb:NN 002123.1
	gb:NM_018487.1 /DEF-Homo sapiens hepatocellular carcinoma-associated animy-miles in 12726 hepatocellular. GEN-HCA112 /PROD-hepatocellular carcinoma-associated animy-miles in 18487.1 (VG=Hs.12126 hepatocellular arcinoma-associated animy-miles in 18487.1
218345_at	carcinoma-associated antigen 112 /FL=gp:ArZassan.i gp:marzan.razan.razan.razar
	Consensus includes gb:AA807056 /FEA=EST /DB_XAKE=g1:2616032 /DB_XAKE=cs:20052235.1 gb:NZ6038.1 gb:NZ6038.1
	dp:195989.1 gb:095819.1 gb:06825.1
	fragment of IgE, high affinity I, receptor 101; alpha polypeption). Desc fragment of IgE, high affinity I, receptorfor; alpha polypeption
211734 g at	move, complete out. (response out.)
	cmo sapiens arginyl-tRNA synthetase (RARS), mRNA. / FEA=mWNA / GEN=FAND. / ENCED=4-55-13-
201330_at	DB_XREF=gi:4506428 /UG=Hs.180832 arginy1-tran synthecase free sint type 7 (FSMA7), mRNA. /FEA-mRNA /
	Homo sapiens proteasome (prosome, macropain) submirt, argument (100 H)
201114_x_at	(prosome, macropain) subunit, alpha type, / / / / / / / / / / / / / / / / / / /
	Consensus includes gD:NM_UVIIS).1 / DEF-MANN September 5 / 100 NOB_XREF=g1:4504514 /UG=Hs.3268 heat shock 70kD protein 6 (HSP70B) / NOB_XREF=g1:4504514 /UG=Hs.3268 heat shock 70kD protein 6 (HSP70B) /
213418_at	FL=gb:NM_002155.1
	gb:AB017493.1 /DEF=Homo sapiens mRNA for DAM-Dinding Zinc Linget (OB) / Company Dinding protein /FL=gb:BC000311/1 propental binding zinc finger(GBF) /DB_XREF=gi:3582142 /UG=Hs.285313 core promoter element binding protein /FL=gb:BC000311/1
208961_s_at	gb:BC004301.1 gb:AP001461.1 gb:AB017493.1 gb:NM_001300.2 mgma /FFB=mRMA /GFN=RPS26 /FROD=riboscmmal protein S26 /
	gb:NM_001029.1 /DEF=Homo sapiens ribosomal protein S26 (MrS20), mayor. /ind.man
21//53_5_ac	gb:NM_004528.1 /DEF=Homo sapiens microsomal glutathione S-transferase 3 (MGST3), mRNA. /FEA=mRNA /GEN=MGST3 / gb:NM_004528.1 /DEF=Homo sapiens microsomal glutathione S-transferase 3 /
202100	PROD=microsomal giucachione 5-Lights crass 3 (2528.1) Prod=microsomal giucachione 5-Lights crass 3 (2528.1) Prod=microsomal giucachione 5-Lights crass 3 (2528.1)
20 CO TO TO	200 related to splicing ractor FRE13 (NULTSO), minuted blicing factor PRP19 /DB_XREF=gi::7657380 /UG=Hs.173980
100000	/GEN=NMP200 / FKUD=nuclear matrix process factor PRP19 /FL=gb:NM_014502.1
203103_S_at	וויריניני איניילי אורייניילי איניילי איניילי איניילי איניילי איניילילי איניילילי

221903 s at	
	Consensus includes gb:AL121934 /DEF-Human DNA sequence from clone RP11-209A2 on chromosome 6. Contains an RPL10 (60S ribosomal protein L10) pseudogene, ESTs, STSs and GSSs /FEA=CDS /DB_XREF=gi:9795199 /UG=Hs.272340 Human DNA sequence from clark contains an RPL10 (60S ribosomal protein L10) pseudogene, ESTs, STSs and GSSs
205120	sapiens CD33 a
100000	gb:NW_G1321.1 /DEF=Homo sapiens U6 snRNA-associated Sm-like protein (LSM4), nRNA. /FEA=mRNA /GEN=LSM4 /FROD=U6 snRNA-associated Sm-like protein /FL=gb:BC000387.1 gb:BC003652.1 gb:AF182290.1 cb. ar.17335.1 cb. NN 012431.1 cb. AF251218.1
201416 at	Consensus includes gb:B6528420 /FEA=EST /DB_XREF=gi:13519957 /DB_XREF=est:602579853F1 /CLONE=IMAGE:4719060 / Fig=Fs. 83484 SRY (sex determining region Y)-box 4 /FL=gb:NM_003107.1
214084 x at	
204861 s at	gb:NN_004536.1 /DEF-Homo sapiens baculoviral IAP repeat-containing 1 (BIRC1), mRNA. /FEA-mRNA /GEN-BIRC1 /PROD-baculoviral IAP repeat-containing 1 /FL=gb:U19251.1 gb:NN_004536.1
221666_s_at	gb:BC004470.1 /DEF=Homo sapiens, clone MGC:10332, mRNA, complete cds. /FEA=mRNA /PROD=Unknown (protein for MGC:10332) / DB_XREF=gi:13325315 /UG=Hs.71869 apoptosis-associated speck-like protein containing a CARD /FL=gb:BC004470.1
218421_at	gb.NM.022766.1 /DEF=Homo sapiens hypothetical protein FLJ23239), mRNA. /FEA=mRNA /GEN=FLJ23239 /PROD=hypothetical protein FLJ23239 /DE,XMED 1223240 /UG=HS.34516 hypothetical protein FLJ23239 /FELJ23239 /DE,XREF=gi:12232440 /UG=HS.34516 hypothetical protein FLJ23239 /FELJ23239 /DE,XREF=gi:1223240 /UG=HS.3440 /UG=HS.34516 hypothetical protein FLJ23239 /FILZ=Gi:10004278 /UG=HS.34516 hypothetical protein FLJ23239 /UG=HS.34516 hypothetical protein FLJ2339 /UG=HS.34516 hypothe
217794 at	sapiens DKFZp564J157 proteir n /DB_XREF=gi:8922156 /UG=Hs.
7 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	gb:NM_003610.1 /DEF-Homo sapiens RAE1 (RNA export 1, S.pombe) homolog (RAE1), mRNA. /FEA=mRNA /GEN=RAE1 / PROD=RAE1 (RNA export 1, S.pombe) homolog /DB_XREF=gi:4506398 /UG=Hs.196209 RAE1 (RNA export 1, S.pombe) homolog / FY=-th-184720.1 ob:NM 003610.1
218055 s at	gb:NW_018268.1 /DEF-Homo sapiens hypothetical protein FLJ10904 (FLJ10904), mRNR. /FER-mRNR /GEN-FLJ10904 / PRON-hypothetical protein FLJ10904 /DB XREF-gi:8922759 /UG-Hs.16470 hypothetical protein FLJ10904 /FL-gb:NW_018268.1
202191 g at	Consensus includes gb:BE439987 /FEA=EST /DB_XREF=gi:9439470 /DB_XREF=est:HTM1-745F /UG=Hs.226133 growth arrest-specific 7 / FT=ch:AB077554.1 gb:RM 005890.1
7	gb:NM 004899.1 /DEF-Homo sapiens brain and reproductive organ-expressed (TNFRSF1A modulator) (BRE), mRNA. /FEA-mRNA / GEN-BNE /PROD-brain and reproductive organ-expressed (TNFRSF1Amodulator) /DE XREF=gi:4757871 /UG=Hs.80426 brain and
203330_S_ac	physical Control of the Control of the Control of the Control of C
217014 of	1840
21,011 at	CONTRACTOR STANGING STANGING AND THEA-EST / DB XREF-gi:2657325 / DB XREF-est:2165b04.sl /CLONE-IMAGE:455119 / INCHES:455119 / CLONE-IMAGE:455119 / INCHES:455119 / INCHES:4551
212386 at	/DB_XREF= fis, clon
18571 s at	protein (HSPC134), mRNA, /FEA-mRNA /GEN-HSPC13 protein /FL-gb:AF212243.1 gb:AF161483.1 gb:NM
10 20 034 000	gb:NM_003751.1 /DEF=Homo sapiens eukaryotic translation initiation factor 3, subunit 9 (eta, 116kD) (EIF3S9), mRNA. / FERA-ENRA /GEN=EIF3S9 /PROD=eukaryotic translation initiation factor 3, subunit 9 (eta, 116kD) /DE_XREF=gi:4503526 / mc-we \$7781 enkaryotic translation initiation factor 3, subunit 9 (eta, 116kD) /FI=gb:U62583.1 gb:NM_003751.1
מיסיים מיסיים	no sapiens hypothetical protein MGC2217 (MGC2217), mRNA.
718047 Sat	T. OO FED TANK OLD THE

	db:NNL 001009.1 / DEF-Homo sapiens ribosomal protein S5 (RRS5), mRNA. /FEA-mRNA /GEN-RPS5 /PROD-ribosomal protein S5 /
	sapiens NADH dehydrogenase (ubiquinone) 1, s FRC2 /PROD=NADH dehydrogenase (ubiquinone) 1, is.193113 NADH dehydrogenase (ubiquinone) 1, s fis.19312 the NADH dehydrogenase (ubiquinone) 1, s
	programment of National Section MGC:2198, mRNA, complete cds. /FEA=mRNA /PROD=Unknown (protein for MGC:2198) / gb:RC000587.1 /DEF=Homo sapiens, clone MGC:2198, mRNA, complete cds. /FEA=mRNA /PROD=Unknown (protein for MGC:2198) / DB_XREF=gi:12653618 /Ug=Hs.227152 mannan-binding lectin serine protease 1 (C4C2 activating component of Ra-reactive
	factor) /FL=gb:BC000587.1 gb:NM_016134.1 /DE=Honoraptens aminopeptidase (LOC51670), mRNA. /FEA=CDS /GEN=LOC51670 /PROD=aminopeptidase /
	DB_XXEE=gg://00000 / 005-nb:2/0979
1	Ildosowar process 22.7 / DEF=Homo Sapiens chromosome 1 open reading frame 7 (CLORF7), mRNA. /FRA=mRNA /GEN=CLORF7 / GD:NM_004895.1 / DEF=Homo Sapiens chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7 / DE_XREF=gi:4757727 /UG=Hs.159483 chromosome 1 open reading frame 7
207075_ac	Fireguing Observations are recommended by Street 1.12920378 /DB_XREF = St.AL567227 /CLONE=CSODF027XA11 (3 prime) / Consensus includes polabeled are recommended by Street 1.12920378 /DB_XREF = St.AL567227 /CLONE=CSODF027XA11 (3 prime) / Fireguing and included by Street 1.12920378 /DB_XREF = St.AL567227 /CLONE=CSODF027XA11 (3 prime) /
201449_ac	Consense includes up in the contract of the co
213720_s_at	SWINK FELGLEU, MALIA BESCHARTON FER FER FOR THE FOR TH
201593_s_at	uncharacterized hypocharanus protein more from the protein kinase (Cam kinase) II gamma (CAMK26), mRNR. / ph.NM_O1222-1, DEF-Home sapies calciumcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / PROP-earliamcalmodulin-dependent protein kinase (CaMkinase) II gamma / DB_XREF_91:4502554 / DB_XREF_91:450
208095_s_at	UG=HS.250857 calciumcalmodulin-dependent protein kinase (CaM kinase) II gamma /KL=gD:U01534.1 gD:NNZ_V04422.1.
	gp:na_cocococc. describution gene C (A.nidulans)homolog /DB_XREF=g1:5729952 /UG=H8.263812 nuclear distribution gene C (A.nidulans)homolog /DB_XREF=g1:5729952 /UG=H8.263812 nuclear distribution gene C (A.nidulans)homolog /Pf=eb:BC002399.1 qb:BC003132.1 gb:AR130736.1 gb:AR125465.1 gb:AR10760.1 gb:NM_006600 I
ZUIII) X at	, mRNA. /FEA=mRNA /GEN=TFG /PROD=TRK-fus
71/839_at	gb:NM_003993.1 / DEF=Homo, sapiens CDC-like kinase 2 (CLK2), transcript variant phclk2, mRNA. / FEA=mRNA / GEN=CLK2 / gb:NM_003993.1 / DEF=Homo, sapiens CDC-like kinase 2 / FI=qb:NM_003993.1 gb:L29218.1
203229_s_at	PROD-CDC-11ke kinase 2 isoloin nelki 100 nelki
203656_at	2033 /UG=Hs.1UU3/ rosapiens syntaxin
218763_at	//FL=gb:AB028/41.1 gb:NM_U10930.1
21384b_ac	CONSENSUS INCLUDES 92:11-12 (DEF-Homo sapiens protein phosphatase 1, regulatory subunit 7 (PPPIR7), mRNA. /FEA-mRNA /GEN-PPPIR7 / PROD-protein phosphatase 1, regulatory subunit 7 /DB_XREF-gi:4506012 /UG-Hs.36587 protein phosphatase 1, regulatory
201214_s_at	subunit 7 /FL=gb:BC000910.1 gb:NM_UNZ/1Z.1
	Consensus includes guitalious / Lie and the gene for ribosomal broken 210, ESTS, GSSs and CpG islands /FEA=mRNA_3 /DB_XREF=gi:8894621 / a gene similar to the gene for ribosomal protein S10, ESTS, STSS, GSSs and chromosome 20 Contains a novel gene, a gene similar to the gene for etherman DNA sequence in STSS, GSSs and CpG islands
216505 X at	gb:NM_022575.1 /DEF=Homo sapiens vacuolar protein sorting 16 (yeast homolog) (VPS16), mRNA. /FEA=mRNA /GEN=VPS16 / PROD=vacuolar protein sorting 16 PROD=vacuolar protein sorting 16
203459_s_at	<pre>// yeast nomolog) / FL=gorn 300001.1 gornm_22.0</pre> gb:NV_004762.1 /DEF=Homo sapiens pleckstrin homology, Sec7 and coiledcoil domains 1(cytohesin 1) (R5CD1), transcript gb:NV_004762.1 /DEF=Homo sapiens pleckstrin homology, variant 1, mRNA. /FRB-PSCD1 /PROD=cytohesin 1, isoform 1 /DB_XREF=gi:4758963 /UG=HS.1050 pleckstrin homology, variant 1, mRNA. /FRB-PSCD1 /FRD-PSCD1 /PROD=cytohesin 1, isoform 1 / DB_XREF=gi:4758963 /UG=HS.1050 pleckstrin homology, variant 1, mRNA. /FRB-PSCD1 /FRD-PSCD1 / PROD=cytohesin 1, isoform 1 / DB_XREF=gi:4758963 / UG=HS.1050 pleckstrin homology, variant 1, mRNA. /FRB-PSCD1 / PROD=cytohesin 1, isoform 1 / DB_XREF=gi:4758963 / UG=HS.1050 pleckstrin homology, variant 1, mRNA. / FRB-PSCD1 / PROD=cytohesin 1, isoform 1 / DB_XREF=gi:4758963 / UG=HS.1050 pleckstrin homology, variant 1, mRNA. / FRB-PSCD1 / PROD=cytohesin 1, isoform 1 / DB_XREF=gi:4758963 / UG=HS.1050 pleckstrin homology, variant 1, mRNA. / FRB-PSCD1 / PROD=cytohesin 1, isoform 1 / DB_XREF=gi:4758963 / UG=HS.1050 pleckstrin homology, variant 1, mRNA. / FRB-PSCD1 / PROD=cytohesin 1, isoform 1 / DB_XREF=gi:4758963 / UG=HS.1050 pleckstrin homology, variant 1, mRNA. / FRB-PSCD1 / PROD=cytohesin 1, isoform 1 / DB_XREF=gi:4758963 / UG=HS.1050 pleckstrin homology, variant 1, mRNA. / FRB-PSCD1 / PROD=cytohesin 1, isoform 1 / DB_XREF=gi:4758963 / UG=HS.1050 pleckstrin homology, variant 1, mRNA. / FRB-PSCD1 / PROD=cytohesin 1 / DB_XREF=gi:4758963 / UG=HS.1050 pleckstrin homology, variant 1, mRNA. / FRB-PSCD1 / PROD=cytohesin 1 / DB_XREF=gi:4758963 / UG=HS.1050 pleckstrin homology, variant 1, mRNA. / GRAPHS-GRAP
202880_s_at	Sec7 and colledcoll domains illeycollesin if the Samonas and Sec7 and colledcoll domains illeycollesin in the Samonas and Sec7 and colledcoll domains illeycollesin in the Samonas and Sec7 and College in the Samonas and Sec7 and

	Cluster Incl. AI201594:qc02h12.xI Homo sapiens cDNA, 3 end /clone=IMAGE-1708487 /clone_end=3 /gb=AI201594 /
	1 H :
ar .	DREPARATE DICTURE TO SEPTIMENT OF THE DESCRIPTION O
	plasma) /FL=gb:UZ45//.1 gp:NM_U05004.1 gp:UZ415/.1 gb.AF020314.1 /DEF=Homo sapiens CMTF-35-H9 mRNA, complete cds. /FEA=mRNA /PROD=CMRF-35-H9 /DB_XREF=gi:4103065 /
s_at.	UG-HS.9688 leukocyte memorane antigen (first) and 1888 leukocyte memorane antigen (first) (fir
	1.1. 1ens McKusick-Kaufman syndrome (MKKS), mRNA. /FEA-mRNA /GEN-MKKS /PROD-McKusi 19055271 /UG-Hs.46743 McKusick-Kaufman syndrome /FL-gb:AF275813.1 gb:AF221992
	gp: AZZIJS3.1 gD: NR. J. JOSEF HOMO Sapiens CGI-39 protein; cell death-regulatory protein GRIM19 (LOC51079), MRNA. /FEA-MRNA / gb: NM 015965.1 / DEF-Homo sapiens CGI-39 protein; cell death-regulatory proteinGRIM19 / DE XREF-gi:7705733 / UG-Hs. 279574 CGI-39 protein; GEN-LOC51079 / PROD-CGI-39 protein; cell death-regulatory proteinGRIM19 / DE XREF-gi:7705733 / UG-Hs. 279574 CGI-39 protein;
220864_s_at	cell death-regulatory protein GKIML9 /rL=go:Aril2370:1 go:Aril23002:1 go:Aril23002:1 GCONE=IMAGE:2499100 /UG=Hs.34578 Consensus includes gb:A1989567 /FEA=EST /DB_XREF=g1:5836448 /DB_XREF=est:ws34e03.x1 /CLONE=IMAGE:2499100 /UG=Hs.34578
213355 <u>at</u>	alpha2,3-sialyltransferase Consensus includes gb.AI809341 /FEA=EST /DB_XREF=gi:5395907 /DB_XREF=est:we96c09.x1 /CLONE=IMAGE:2348944 /UG=Hs.170121
212587_s_at	protein tyrosine phosphatase, receptor type, the protein the protein type of the protein type of the protein to
2027278_ac	DE ANGE - 9:10031128 / 702 MENT - 1 MENT. / gb:NM 004699.1 / DEF-Homo sapiens DNA segment on chromosome X (unique) 9928 PEA-MENT. / PEA-MENT. / FEA-MENT. / GEN-DXS928E / Unique) 9928 PEA-MENT. / GEN-DXS928E / TOCAMOSOME X (unique) 9928 PEA-MENT. / GEN-DXS928E / TOCAMOSOME X (unique) 9928
203262_s_at	expressed sequence /klaga:Brunouzo.1 gb:moszco.1 gb:moszco.1 gb:moszco.2 consensus /CLONE-ADAAGD10 / Consensus includes gb:aVV0118 /FBE-EST /DB_XREF=gi:10717648 /DB_XREF=est:AV701318 /CLONE-ADAAGD10 /
213357_at	UN=HS./10820 ArgADJ-INCERACTING PIOCEAN SEPTEMBER (HASJ4442), MRNA. /FEA-MRNA /GEN=HASJ4442 / DEN CONTROL OF THE CONTROL OF TH
מייים בייים	gb:NW_GO04231.1 /DEF=Homo sapiens AFPase, vacuolar, 14 kD (ATP6S14), mRNA. /FEA=mRNA /GEN=ATP6S14 /FROD=ATPase, vacuolar, 14 kD /FL=gb:D49400.1 gb:NM_G04231.1
20132/_at	dehydrogenase, type II (HADH2), mRNA. /FEA-mRNA /GEN-HADH REF=g1:4758503 /UG-HS.171280 hydroxyacyl-Coenzyme A dehy 96132.1 gb:aF035555.1 gb:AF069134.1 gb:NM_004493.1
213735_s_at	Consensus includes gb:AI557312 /FEA=EST /DB_XREF=gi:4489675 /DB_XREF=est:PT2.1_16_B11.r /UG=Hs.1342 cytochrome c oxidate subunit Vb
212472_at	Consensus includes gb:BE965029 /PEA-EST /DB XREF=g1:11/089/0 /DB_XREF=est:0010001212 /CCAL-1710239 /UG=HS.239483 Human clqne
214805_at	Consensus includes go: 07273.1 / Diff. Linear Containing ion transport regulator 5 (FXYD5), mRNA. /FEA=mRNA /GEN=FXYD5 / Gb: NM_014164.2 / DEF=Homo sapiens FXYD domain-containing ion transport regulator 5 / Containing ion transport regulator 5 / Contai
218084_x_at	PROD=related to ion channel /DB ArkF gi:lioizouw / UG-in: 2015 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
	<pre>gb:NM_024513.1 /DEF=Homo sapiens FYVE and colled-coll domain containing 1 / PROPERTYPE and colled-coll domain containing 1 / DB_XREF=gi:13470091 / UG=HS.257267 PYVE and colled-coll domain containing 1 / PROPERTYPE and colled-coll domain containing 1 / PROPERTYPE and colled-coll domain containing 1 / PROPERTYPE PROPERTYPE and colled-coll domain containing 1 / PROPERTYPE AND 124513.1</pre>
22-0-0-1	

	gb:NM_000992.1./DEF=Homo sapiens ribosomal protein L29 (RPL29), mRNA. /FEA=mRNA (GEN=RPL29 / PROD=ribosomal protein L29 /
200823_x_at	DB_XREF=gi:4506628 /UG=Hs.183698 Illoscural protein, close MGC:2478, mRNA, complete cds. /FEA=mRNA /PROD=hypothetical gi:2408, mRNA, complete cds. /FEA=mRNA /PROD=hypothetical protein, close MGC:2478, mRNA, complete cds. /FEA=mRNA /PROD=hypothetical protein, close MGC:2478, mRNA, complete cds. /FEA=mRNA /PROD=hypothetical
208968_s_at	protein /DB_XRRF=gi:12803484 /UG=Hs.4900 hypothetical protein /FL=go:AF40504:1 gy:CCC051 gy ARD / Clone MGC:2053, mRNA,
	<pre>gb:BC000733.1 /DEF=Homo sapiens, eukaryotic translation initiation factor 3, subunit 4 (delta, 44kD) /DE_XREF=gi:12653882 / complete cds. /FEA=mRNA /PROD=eukaryotic translation initiation factor 3, subunit 4 (delta, 44kD) /FL=gb:AF094850.1 gb:BC000733.1 gb:AF020833.1 UG=Hs.28081 eukaryotic translation initiation factor 3, subunit 4 (delta, 44kD) /FL=gb:AF094850.1 gb:BC000733.1 gb:AF020833.1</pre>
208887_at	gb:U96074.1 gb:NM_003755.1 /PEA=EST /DB_XREF=g1:5877913 /DB_XREF=est:wv03e06.x1 /CLONE=IMAGE:2528482 /UG=HS.1948 ribospmal
214097_at	protein 521 /CLONE=IMAGE: 3057859. /
212191 x at	Consensus includes gb:AW574664 /FEA=EST / Lb_AKEF=91:/240203 / Lb_AKF=91:/240203 /
	opiens similar to mouse Glt3 or D. malanogaster transcription ractor in 580 /PROD-similar to mouse Glt3 or D. malanogastertranscription factor II 2.79818 similar to mouse Glt3 or D. malanogaster transcription factor IIE
217957_at	/FL=gb:BC005152.1 gb:AF093680.1 gb:MM_U12442.1 Comesses: includes db:AU151560 /FEA=EST /DB_XREF=g1:11013081 /DB_XREF=est:AU151560 /CLONE=NT2RP2005555 /UG=Hs.242894
201658_at	ADP-ribosylation factor-like 1 /FL-gb:NM 001177.2 gb:L28997.1
	rame 59 (CZIOKE55), ukwa: -Hs.5811 chromosome 21 open 1
218123_at	/FL=gb:iM 031254.1 gp:BCU0/19:1 gp:RAL_OLIDSTILE_STATE GP:RESTATE (reast) homolog 2 (SCO2), nuclear gene
	gb:NN 005138.1 / DEF-Homo Saptens Sto (cytochtome oxidase deficient, yeast) encoding mitochondral product nrn-us, 27813 Sto (cytochtome oxidase deficient, yeast) homolog 2 /FL-gb:NN 005138.1
205241_at	nomolog 2 / DB AKER 9: 4000521 / OG 115: 100 PROSPERS (MPHOSPH6), MRNA. /FEA-MRNA /GEN-MPHOSPH6
203740_at	/PROD=M-phase phosphoprotein 6 /DB XREF=gi:5031918 /UG=HS.152/20 M-phase phosphoprotein 6 /LF-g-man / GEN=MGC3133
221263 s at	gb:NM 031287.1 / DEF=Homo Sapieus influencearea processis 13775199 /FL=gb:NM 031287.1
	: fragment of IgG, receptor, transporter, alpha ("Carri,") hent of IgG, receptor, transporter, alpha (DB_XREF=gl:475) hent of IgG, receptor, slabs (FF=ch:NM 004107:1 db:U1225)
218831_s_at	
	06196 8.1
201400_at	NOGERS. 02.73 PINICASCONE PERCEPTOR ACCEPTOR [(PREMACE), MRNA. /FERA-MRNA /GEN-RABACI /PROD-Rab acceptor 1
203136_at	(prenylated) /DB_XREF=gi:5453959 /UG=HB.11417 Rab acceptor 1 (prenylated) /FEA=mRNA /GEN=DF
205382 s.at	597 D component of
,	ogen
208/14_ac	Liavoprocent 1 (510) 12 52. NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 2 (8kD, B8), gb:BC003674.1 /DEF=Homo sapiens, NADH dehydrogenase (ubiquinone) 1 alphasubcomplex, 2 (8kD, B8) clone MgC:12315, mRNA, complete cds. /FEA=mRNA /FROD=NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 2 (8kD, B8)
209224_s_at	7029.1
210574_s_at	gb:AF241788.1 /DEF=Homo sapiens NPD011 (NFD011) MANA, Compress homolog /FL=gb:AF241788.1 /UDE-Homo sapiens NPD011 nuclear distribution or present of the contraction
214259 s_at	Consensus includes gb:AI144075 /FEA=EST /DB_XREF=91:3605884 /DB_XREF=StrigioSez /UG=Hs.6980 aldo-keto reductase family 7, member A2 (aflatoxin aldehyde reductase)

209234_at	
203654 s at	gb:NML_004645.1 /DEF-Homo sapiens coilin (COIL), mRNA. /FEA-mRNA /GEN-COIL /PROD-coilin /DB_XREF-gi:4758023 /UG=Hs.966 coilin / FL-gb:U06632.1 gb:NM_004645.1
201129 at	
10000	gb:EC000365.1 /DEF=Homo sapiens, general transcription factor IIH, polypeptide 1 (62kD subunit), clone MGC:8323, mRNA, complete cds. /FEA-mRNA /PROD=general transcription factor IIH, polypeptide 1(62kD subunit) /DB_XREF=gi:12653194 (GG-HS 98578 general transcription factor IIH, polypeptide 1 (62kD subunit) /FLeah:RC000365.1 db:M95809.1 db:NM,005316.1
218072 at	այտ
203063_at	gb:RM_014634.1 /DEF-HGmo sapiens KIBA0015 gene product (KIBA0015), mRNA. /FEA-mRNA /GEN-KIBA0015 /PROD=KIBA0015 gene product /DB_XREF-g1:7661861 /UG-HS.278441 KIBA0015 gene product /FL-gb:D13640.1 gb:NM_014634.1
204037 at	Consensus includes gb:BF055366 /FEA-EST /DB_XREF=gi:10809262 /DB_XREF=est:7j78f10.x1 /CLONE=IMAGE:3392587 /UG-Hs.75794 endothelial differentiation, lysophosphatidic acid G-protein-coupled receptor, 2 /VE-Gb:U78192.1 gb:U80811.1 gb:UM_001401.1
218298 s at	
204158 s at	gb:NM 006019.1 /DEF-Efono sapiens T-cell, immune regulator 1 (TCIRGI), mRNR. /FEA-mRNA /GEN-TCIRGI /PROD-ATPase, H+ transporting, 116kD /DB XREF-cq:5174620 /UG-Hs.46465 T-cell, immune regulator 1 /FL-gb:U45285.1 gb:NM 006019.1
208842 s at	787 /FEA=EST /DB_XREF=gi:1422930 /DB_XREF=est:zd9
203047 at	pp.NNC_005990.1 / DEF-Homo sapiens serinethreonine kinase 10 (STK10), mRNA. /FEM-mRNA /GEN=STK10 / PROD=serinethreonine kinase 10 / DB XREF=cd:5174700 /UG=Hs.16134 serinethreonine kinase 10 /FL=gb:AB015718.1 gb:NM_005990.1 gb:AF119894.1
14.08 0.01	
	gb:BC002877.1 /DEF=Homo sapiens, Similar to hypothetical protein FLJ11585, clone MGC:11258, mRNA, complete cds. /FEA=mRNA / PROD=Similar to hypothetical protein FLJ11585 /DB_XREF=gi:12804048 /UG=Hs.315367 Homo sapiens, Similar to hypothetical protein
209858_x_at	FLJ11585, Clone McC:11258, mkWA, complete cas /rlegu:bcock// Cluster Incl. U30894:Human N-sulphoglucosamine sulphohydrolase mRWA, complete cds /cds=(12,1520) /gb=U30894 /gi=1173542 /
35626_at	ug=Hs.31074 /len=2657 gb:NM_015853.1 /DEF=Homo sapiens ORF (LOC31035), mRNA. /FEA=mRNA /GEN=LOC51035 /PROD=unknown protein LOC51035 /
2018/1_s_at	nm clone DKFZp564G20 is.16492 DKFZP564G20
100000000000000000000000000000000000000	gb:NW_022731.1 /DEF=Homo sapiens similar to rat nuclear ubiquitous casein kinase 2 (NUCKS), mRNA. /FEA=mRNA /GEN=NUCKS / PROD=similar to rat nuclear ubiquitous casein kinase2 /DB_XREF=gi:12232386
201960 s at	sapiens KIAA0916 protein (KIAA) 151411 KIAA0916 protein /FL=g
47608_at	Cluster Incl. AI697401:tq18h09.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2209217 /clone_end=3 /gb=A1697401 /gi=4985301 / ug=Hs.9741 /len=639
210250_x_at	gb.AF067854.1 /DEF=Homo sapiens adenylosuccinate lyase (ADSL) mRNA, alternatively spliced, complete cds. /PFS-APS-APS-HOMO (GEN=ADSL /PROD=adenylosuccinate lyase /PL=gb:AF067854.1
218495_at	gb.NM_004182.1 / DEF=Homo sapiens ubiquitously-expressed transcript (UXT), mRNA. /FEA=mRNA /GEN=UXI /PROD=ubiquitously-expressed transcript / DB_XRRF=gi:4759297 / UG=Hs.172791 ubiquitously-expressed transcript //FL=gb:BC000720.1 gb:AF092737.1 gb:NM_004182.1 gb:AF083241.1 gb:AF083242.1

	3_XREF=gi:4504106 /UG=Hs.2706 glutathione p
701100 ac	print Oction (PER-Homo sapiens chromosome X open reading frame 5 (CXORFS), mRNA. /FEA-mRNA /GEN-CXORFS / Grant Oction 1 gene /FL-gb:NM_003611.1 DEF-Homo sapiens chromosome X open reading from fast oral-facial-digital syndrome 1 gene /FL-gb:NM_003611.1
203569_s_at	.523904 /FEA=EST /DB_XREF=g1:1278739
201316_at	UG-HS.181309 proteasome (prosome, macropain) subunit, alpha type, / /ri-go:nm.ouz/e/
218336_at	gp:nm_012394.2 /DEF-main Bagrens 2 (201394.2 gb:AF165883.1 gb:AF117237.1 gb:AF151065.1 UG-HS.298229 prefoldin 2 /FL-gb:NM_012394.2 gb:AF165883.1 gb:AF117237.1 gb:AF151005.1 UG-HS.198100 /UG-HS.167679
204370 s at	Consensus includes gb: BE502377 /FEA=EST /DB_XREF==51:9/04/85 /DB_XREF==51:1942903.x1 /CDGANINCLUDENT /CDGANINCLUDEN
	gb:NM_005327.1 /DEF=Homo sapiens L-3-hydroxyacyl-Coenzyme A dehydrogenase, short chain (HADHSC), mWNA. /FEA=mwNa /vEm=nn45c / PROD=L-3-hydroxyacyl-Coenzyme A dehydrogenase, shortchain /DB_XREF=gi:4885386 /UG=Hs.8110 L-3-hydroxyacyl-Coenzyme A dehydrogenase,
201036_s_at	short chain /PI-gb: BC000306.1 gb:NM_U03347.1.
208776_at	(prosome, macropain) 26S subunit, non-ATPase, 11 /FL=gb:BC000437.1 gb:BC004430.1 gb:AB003102.1 gb:AFVU1212.1 gb:ARVU1212.1 gb:AR
214765_s_at	
204225 at	gb:NM_006037.2 /DEF=Homo sapiens histone deacetylase 4 (HDAC4), mRNA. /FEA=mRNA /GENHDAL4 /FROD-miscome wearest
25-03-03	Consensus includes gb:AA669797 /FEA=EST /DB_XREF=g1:2631296 /DB_XREF=est:ag36c01.s1 /CLONE=IMAGE:1118880 /UG=HS:/DGJ3
214170_x_at	/FEA=mRNA /GEN=FLJ12171
218210 at	protein FL01211 / DB XRRF egi: 13375839 / UG=Hs.31431 hypothetical protein FL012171 / FL=Gp: AL135631.1 ginn _ UGABL: 1 protein FL012111 / DB XRRF egi: 13375839 / UG=Hs.31431 hypothetical protein FL012111 / DB XRRF egi: 13375839 / UG=Hs.31431 hypothetical protein FL012171 / FL=Gp: Al135631.1 ginn _ UGABL
	gb:NM_017733.1 /DEF=Homo sapiens hypothetical protein FLJ20265 (FLJ20265), mRNA. /FEA=mRNA /GEN=FLJ20265 /FKDJ=LJ20265 /FEA=mRNA /GEN=FLJ20265 /FEA=MR
218652_s_at	process Florago / Do Andr - 9. Construction of the control of the
201270_x_at	DB_XREF=gi:13357209 /UG=Hs.4770 KIAA1068 procein /FL-gi:ccoopsile gi:13357209 /UG=Hs.4764 KIAA0763 gene
203906_at	Consensus includes gb:Aloszews (Francis) (Fran
	gb: BC002587.1 /DEF-Homo sapiens, hypothetical protein, clone MGC:1067, mKNA, complete cds. /rea-mann /race-ing protein /FL-gb: BC002587.1
15-0-10-10-10-10-10-10-10-10-10-10-10-10-1	gb: BC003679.1 /DEF-Homo sapiens, ATP synthase, H+ transporting, mitochondrial FO complex, subunit e, clone MGC:12532, mkWW,
209492_x_at	UG-HS. 85539 ATP Synthase, H+ transporting, mitochondrial FO complex, subunit e /FL=gb:BC0036/9.1
7e 2 800010	Consensus includes gb:AM245400 /FEA-EST /DB_XREF=G1:5588393 /DB_AXGF=G8t:2022/J1:JLIME /VDCM2-AM245400 /FEA-EST /DB_AXGF=G1:5588393 /DB_AXGF=G8t:2022/J1:JLIME /VDCM2-AM245400 /FEA-EST /DB_AXGF=G1:5588393 /DB_AXGF=G8t:2022/J1:JLIME /VDCM2-AM245400 /FEA-EST /DB_AXGF=G1:5588393 /DB_AXGF=G8t:2022/J1:JLIME /VDCM2-AM245400 /FEA-EST /DB_AXGF=G8t:2022/J1:JLIME /VDCM2-AM245400 /FEA-EST /DB_AXGF=G8t:2022/JI:JLIME /VDCM2-AM245400 /FEA-EST /DB_AXGF
22-0-0-0-1-1	gb:NM_006834.1 /DEF=Homo sapiens RAB32, member RAS oncogene family (RAB32), mRNA. /FEA=mRNA /GEN=RAB32 /FROD=RAB32, member RAS gb:NM_006834.1 gb:NM_006834.1
204214_s_at	thetical protein FLJ20657 (FL
217796_s_at	protein FL720657 /DB_XREF=g1:8923608 /UG-Hs.164256 hypothetical protein FL720657 /FL-gp:NM_UL/241.1
Ťe 705500	Consensus includes gb:NZ2903 /FEA=EST /DB_XREF=g1:113/U53 /DB_XREF=eSC:YX00eU4:51 /CLCAL-LINGEST /DB_XREF=EST:113/U53 /DB_XREF=eST:YX00eU4:51 /CLCAL-LINGEST /DB_XREF=EST:YX00eU4:71 /CLCAL-LINGEST /D
227.22000	gb:AF034607.1 /DEF-Homo sapiens chloride channel ABF mRNA, complete cds. /FEA-mRNA /PROD=chloride channel ABF /UD_AREF-94:1940000 / PROD=chloride channel ABF /UD_AREF-94:194000 / PROD=chloride intracellular channel 1 /FL-qb:U93205.1 gb:AF034607.1 gb:AF109197.1 gb:NX_001288.2
200033_ac	Consensus includes gb:AJ400843.1 /DEF=Homo sapiens partial mRNA for imminoglobulin-like cell surface receptor FDFU3-m14, Solution Consensus includes gb:AJ400843.1 /DEF=Homo sapiens partial mRNA for imminoglobulin-like cell surface receptor FDFU3-m14, Solution Consensus includes gb:AJ400843.1 /DEF=Homo sapiens partial mRNA for imminoglobulin-like cell surface receptor FDFU3-m14, Solution Consensus includes gb:AJ400843.1 /DEF=Homo sapiens partial mRNA for imminoglobulin-like cell surface receptor FDFU3-m14, Solution Consensus includes gb:AJ400843.1 /DEF=Homo sapiens partial mRNA for imminoglobulin-like cell surface receptor FDFU3-m14, Solution Consensus includes gb:AJ400843.1 /DEF=Homo sapiens partial mRNA for imminoglobulin-like cell surface receptor FDFU3-m14, Solution Consensus includes gb:AJ400843.1 /DEF=Homo sapiens graph consensus includes gb:AJ400843.1 /DEF=Homo sapiens graph consensus includes gb:AJ400843.1 /DEF=Homo sapiens graph consensus graph consens
222218_s_at	alternative torm. /FEASTHANA /GENSFUROS-Bit / FAUD-CELL Buriana and Anna Anna Anna Anna Anna Anna An
1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	gb:AF015451.1 /DEF=Homo sapiens Usurpin-beta mRNA, complete cds. /FEA=CDS /FROD=Usurpin-beta /DLL
211862_X_at	

, 08000C	gb.aF118652.1 /DEF-Homo sapiens PKCq-interacting protein PICOT (PICOT) mRNR, complete cds. /FEA-mRNR /GEN-PICOT /PROD-PKCq- interacting profein PICOT /DB XREF-c1:6840952 /UG-Hs.42644 thioredoxin-like /FI-gb:BC005289.1 gb:AF118649.1 gb:AF118652.1
	gb:NM_001398.1 /DEF=Homo sapiens enoyl Coenzyme A hydratase 1, peroxisomal (ECH1), mRNA. /FEA=mRNA /GEN=ECH1 /PROD-peroxisomal enoyl-coenzyme A hydratase 1, peroxisomal /
	FLEGD:NM_U01398.1 gp:U10000.1 Cluster Incl. AIG5857:wb65b10.x1 Homo sapiens CDNA, 3 end /clone=IMAGE-2310523 /clone_end=3 /gb=AI654857 /gi=4738836 /
	ugens. 1001147.1 /DEF-Homo sapiens ectonucleoside triphosphate diphosphohydrolase 6 (putative function) (ENTPD6), mRNA. / gb.NM_001247.1 /DEF-Homo sapiens ectonucleoside triphosphate diphosphohydrolase 6 (putative function) /DB_XREF-gi.4557422 / FEA-mRNA /GEN-ENTPD6 /PROD-ectonucleoside triphosphate diphosphohydrolase 6 (putative function) /DB_XREF-gi.4557422 /
	UG-HS.1230 ectonucleoside triphosphate diphosphonydronase o the control of the co
718026_ac	DB_XREF=girroring for marked to the control of the
2141/1-X-at	Consensus includes 9:4L031714 (DEF-Human DNA sequence from clone LA16-358B7 on chromosome 16 Contains the UBE21 gene for Consensus includes 9:4L031714 (Def-Human DNA sequence from Clone LA16-358B7) and an RPS20 (40S Ribosomal protein S20) pseudogene. Contains ESTs.
	STSS. GSSs and a putative CpG is /FEA=mRNA /DB_XREF=gi:4775608 /UG=HS.84285 ubiquitin-conjugating enzyme E21 (nometagous to yeast UBC9) /FL=gb:U45328.1 gb:U31933.1 gb:BC000427.1 gb:U31882.1 gb:U31882.1 gb:U66818.1 gb:U66867.1 gb:U38785.1 gb:U345.1 gb:U39092.1
212607 at	ncludes gb:N32526 /FEA=EST /DB_XREF=gi:1152925 /DB_XREF=est:yy11f04.s1 /CLONE=IMAGB: on cancer antiqen 8
. 64418 at	Cluster Incl. AI472320:tj87c02.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2148482 /clone_end=3 /gb=AI472320 /gi=4334410 /
11 00 00 00 00 00 00 00 00 00 00 00 00 0	9-3-3-4-3-4-4-4-4-4-4-4-4-4-4-4-4-4-4-4-
218019 s at	
219220 x at	gb.NM 020191.1 /DEF=Homo sapiens GK002 protein (GK002), mRNA. /FEA-mRNA /GEN=GK002 /PROD=GK002 protein /DB_XMEF=g1:9910243 / UG-HS.107127 GK002 protein; gibt protein; chromosome 3 open reading frame 5 /FL=gb:AF321613.1 gb:AF328645.1 gb:NM_020191.1
212355_at	Consensus includes gb:A1075450 /FEA=EST /DB_XREF=gi:3399805 /DB_XREF=est:oz82g10.x1 /CLONE=IMAGE:1681890./ UG=Hs.7911 KIAA0323 protein
214735_at	166711 /FEA=EST /DB_XREF=qi:6398236 /DB_XR otein
219041_s_at	o sapiens zinc finger protein (AP4), mRNA. /FEA=mRNA /GEN=AA4 /PKOD=Zinc Ling 18.90693 zinc finger protein /FL=gb:BC000363.1 gb:NM 014374.1
209110 s_at	gb:AL050259.1 /DEF=Homo sapiens mRNA; cDNA DKFZp564D0782 (from clone DKFZp564D0782); complete cds. /rea=mNA /sex=nNA_004761.1 PROD=hypothetical protein /DB_XREF=gi:4886476 /UG=Hs.170160 RAB2, member RAS oncogene family-like /FL=gb:D85757.1 gb:NA_004761.1 gb:AL050259.1
215691_x_at	Consensus, includes gb:AV702994 /FEA=EST /DB_XREF=g1:10719324 /DB_XREF=est:AV702994 /CLONE=ADBAPB06 /UG=HS.46967 HSFC034 protein
207614_s_at.	gb:NM_003592.1 /DEF=Homo sapiens cullin 1 (CUL1), mRNA. /FEA=mRNA /GEN=CUL1 /PROD=cullin 1 /DB_KKEF=g1:4505100 / UG=H8:14541 cullin 1 /FL=gb:U58087.1 gb:NM_003592.1
213405_at	Consensus includes gb:N95443 /FEA=EST /DB_XREF=gi:1267753 /DB_XREF=est:2D81cil.s1 /CLUNE=1MA6E:31UUU0 /UG-ms:17100 MUMA
208835 <u>.s_at</u>	Consensus includes gb:AM089673 /FEA=EST /DB_XREF=g1:604/01/ /DB_XREF=est:xdinil.xi /CLOMB-IRAND:207100 /CCLOMB-IRAND:207100 /CCLOMB-IRAND:207100 /CCLOMB-IRAND:207100 /CCLOMB-IRAND:207100
202225_at	XX / CLOND - LEGGE & COLOR - C
218358_at	gb:NM_024324.1 /DEF=Homo sapiens hypothetical protein MGC11256 (MGC11256), mRNA. /FEA=mRNA /GEN=MGC11256 /PROD=hypothetical protein

	MGC11256 /DB_XREF=gi:13236568 /UG=Hs.28029 hypothetical protein MGC11256 /FL=gb:BC002894.1 gb:NM_024324.1
	s gb:Al167164 /FEA=EST /DB_XREF=g1:3700334 /DB_XREF=est:0008c10.x1 /CLONE=INAGE:1565586 /UG=Hs.23200
213511_s_at	related protein 1 gb:NM_021824.1 /DEF=Homo sapiens NIF3 (Ngg1 interacting factor 3, S.pombe homolog)-like 1 (NIF3L1), nRNA. /FEA=mRNA /GEN=NIF3L1 / gb:NM_021824.1 /DEF=Homo sapiens NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:11141898 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:11141898 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:11141898 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:11141898 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:1114188 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DE_XREF=gi:111418 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombehomolog)
218133_s_at	S.pombe homolog)-like 1 /FL=gb:AF182416.1 gb:NM_UZI844.1 gb:AFUBLISH: \$\frac{1}{2}\text{SER} \text{SER}
212007_at	domain-containing 1 GD:NM_004853.1 /DEF=Homo sapiens syntaxin 8 (STX8), mRNA. /FEA=mRNA /GEN=STX8 /PROD=syntaxin 8 /DB_XREF=g1:4759187 /UG-Hs.119525
204690_at	syntaxin 8 /FL=gb:AF062077.1 gb:AF016.15.1 gb:AF113323.1 gb:MR_complete cds. /FEA=mRNA /PROD=cathepsin X precursor / gb:AF073890.1 /DEF=Homo sapiens cathepsin X precursor, mRNA, complete cds. /FEA=mRNA /PROD=cathepsin X precursor /
210042_s_at	DB_XREF=g1:3650437 /UG=HS.252549 cathepsin z /rin=pirroffsin complete cds /cds=(3531,4286) /gp=AB007915 /gi=3413853 / Cluster Incl. AB007915:Homo sapiens mRNA for KIAA0446 protein, complete cds /cds=(3531,4286) /gp=AB007915 /gi=3413853 /
76070	no sapiens encyl Coenzyme A hydratase, short chain, 1; mitochondrial (ECHSI), nuclear gene encoding mRNA. /FEA-mRNA /GEN-ECHSI /PROD-mitochondrial short-chain encyl-coenzyme Ahydratase 1 precursor / new 16784 encyl Coenzyme A hydratase, short chain, 1, mitochondrial /FL-gb:NM_004092.2 gb:D13900.1
201135 ac	DE ARLT-91:12:0757 CO. S. PROD-hypothetical protein PLJ10024 (FLJ10024), mRNA. /FEA-mRNA /GEN-FLJ10024 /FROD-hypothetical protein PLJ10024 /FR-HOME STREE-GING OF STREE-GI
28 8 28 28 28 28 28 28 28 28 28 28 28 28	gb:NM 002198.1 /DEF=Homo sapiens interferon regulatory factor 1 (IRF1), mRNA. /FEA=mRNA /GEN=IRF1 /PROD=interferon regulatory factor 1 /FL=qb:NM 002198.1
202531_at	XREF=g1:12019514 /DI
213465_s_at	protein phosphatase 1, requarcity submitted a acyltransferase 1 (peroxisomal 3-oxoacyl-Coenzyme A thiolase) (ACAAI), girlw. 01607.2 / DEF=Homo sapiens acktyl-Coenzyme A acyltransferase 1 / defined and acyltransferase 1 / defined a
202025 x at	s.166160 acetyl-Coenzyme A acylt: 01607.2
203944_x_at	gb:NR_007049.1 /DEF=Homo sapiens butyrophilin, subfamily 2, member Al (BTN2Al), mRNA. /FEA=mRNA /GAN-DIMAL. JAN-DIMAL JAN-
214290_s_at	Consensus includes gb:A131334 /FEA=EST /DB_XREF=g1:4018929 /DE_ARGE=est:La//lux.az /caca /CEA_=ING1 /PROD=inhibitor of family, member 0
208415_x_at	gb:NM_005537.1 / DEF=Homo sapiens inhibitor of growth 1 family, member 1 (Lamily, member 1 / DB XREF=gi:5031792 / UG-Hs.46700 inhibitor of growth 1 family, member 1 / DB XREF=gi:5031792 / UG-Hs.46700 inhibitor of growth 1 family, member 1 / DB XREF=gi:4406694 / UG-Hs.286027
216396_s_at	Consensus includes gb:AF131850.1 / DEF-Homo sapiens clone 44900 mans Sequence. / Consensus includes gb:AF131850.1 / DEF-Homo sapiens clone 44900 mans / PER-HOND-fumarate hydratase / DB_XREF-gi:4503716 / etcposide-induced mans
203033 x_at	000143.1 (SLC31A1), mRNA. /FEA=mRNA /
203971 at	gb:NW_001859.1 /DEF=Homo sapiens solute darrier lamily -1 (Logberm.spr.com) / 1/DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family GRN=SLC31A1 /PROD=solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /DB_XREF=g1:4507014 /UG=Hs.73614
203241 at	gb:NM_003369.1 /DEF=Homo sapiens UV radiation resistance associated gene (UVRAG), mRNA. /FEA=mRNA (GEN=UVRNA /FNA) / LANG 003369.1 FNA) FN
201142_at	Consensus includes gb:AA577698 /FEA=EST /DB_XREF=g1:2355882 /DB_XREF=est:nn22AN0.sr /CLONN=Intac:10.0009 /OS. translation initiation factor 2, subunit 1 (alpha, 35kD) /FL=gb:E0002513.1 gb:Nn /Od0094.1
208818 s.at	Vicransierase, cione 1 653300 /UG=Hs.240013
203436_at	gb:NM_006413.1 /DEF=Homo sapiens ribonuclease P (30kD) (RPP30), mRNA, /FEA=mRNA /GEN=RFF30 /ENCJ=LECTION / FEA=mRNA / GEN=RFF30 / ENCJ=LECTION / FEA=mRNA / GEN=RFF30 / ENCJ=LECTION / FEA=mRNA / GEN=RFFF30 / ENCJ=LECTION / ENCJ=LECTION / FEA=mRNA / GEN=RFFF30 / ENCJ=LECTION /

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sapiens cartilage associated protein (CKTAP), mRNA. /FEA-mRNA /GEN-CRT.1600 /UG-Hs.155481 cartilage associated protein /FL-gb:NM_006371.1 sapiens KIAA0266 gene product (KIAA0266), mRNA. /FEA-mRNA /GEN-KIAA026 Hs.127376 KIAA0266 gene product /FL-gb:NM_021645.1 gb:D87455.1
o sapiens KIAA0266 gene product (KIAA0266), mRNA. /FEA-mRNA /GEN-KIAA0266 /PROD-KIAA0266 Hs.127376 KIAA0266 gene product /FL-gb:NM_021645.1 gb:D87455.1.
gb.AF100752.1 /DEF-Homo sapiens transitional endoplasmic reticulum AFPase mRNA, complete cds. /FEA-mRNA /PROD-transitional and and an analysis of the second of the second and an arrangement of the second and an arrangement of the second and arrangement of the second and arrangement of the second and arrangement of the second arr
endopiasame retroriam mires (12 per 17 pe Consensus includes gb:AV18192 (FEA=EST / DE XEEF=gi:10815344 / DE XEEF=est:AV718192 / CLONE=FHTAABE08 / UG=Hs.171957 triple functional domain (FFFF increacing) /FL-qb:RF091395.1
Consensus includes gb:293241 /DEF=Human DNA sequence from clone 222E13 on chromosome 22. Contains three novel genes, an ATP Synthase G Chain, Mitochondrial (EC 3.6.1.34) pseudogene and the DIAl gene for diaphorase (NADH) (cytochrome b-5 reductase) (FC 1.6.2.2). Contains ESTS. STSS /FEA-mRNA 5 /DB XRRF-gi:4826450 /UG-HS.278314 hypothetical protein
Consensus includes gb:AB023216.1 / DEF-EGomo saptens mRNA for KIAA0999 protein, partial cds. /FEA-mRNA /GEN-KIAA0999 / RPROD-KIAA0999 protein /DB XREF-gi:4589641 /UG-Hs.4278 KIAA0999 protein
Consensus includes gb.AI819709 /FEA=EST /DB_XREF=gi:5438788 /DB_XREF=est:wj77c07.xl /CLONE=IMAGE:2408844 /UG=Hs.108104 ubiquitin-conjugating enzyme E2L 3 /FL=gb:NM_003347.1
Consensus includes gb:A1123233 /FEA=EST /DB_XREF=g1:3538999 /DB_XREF=est:qa47e04.xl /CLONE=IMAGE:1689918 /UG=Hs.167496 RAN binding protein 6
gb:NW_017646.1 /DEF=Homo sapiens tRNA isopentenylpyrophosphate transferase (IPT), mRNA. /FEA=mRNA /GEN=IPT / PROD=FRNA isopentenylpyrophosphate transferase /DB_XREF=gi:8923064 /UG=Hs.288036 tRNA isopentenylpyrophosphate transferase / FL=qb:NM 017646.1
EE-Homo sapiens glucose phosphate isomerase (GPI), mRNA. /FEA-mRNA /GEN=GPI /FROD=gluco 6 /UG-Hs.180532 glucose phosphate isomerase /FL-gb:BC004982.1 gb:K03515.1 gb:NM_000175.
gb:AF112207.1 /DEF-Homo sapiens translation initiation factor eIF-2b delta subunit mRNA, complete cds. /FEA-mRNA / PROD-translation initiation factor eIF-2b deltasubunit /DB_XREF-g1:6563201 /UG-Hs.169474 DKFZE586J0119 protein /
programmer of the property of the protein mRNA, complete cds. /FEA=mRNA /PROD=KIAA0220-like protein / gp:aF2220691 / DEP=Homo sapiens KIAA0220-like protein mRNA, complete cds. /FE-gb:AP229069.1 PR XFRE-d:891670 /HG=Hs. 251928 nuclear pore complex interacting protein /FL-gb:AP229069.1
Consensus includes gb:AI184802 /FEA=EST /DB_XREF=gi:3735440 /DB_XREF=est:qd24g04.x1 /CLONE=IMAGE:1724694 /UG=Hs.8551 PRP4SfKWD splicing factor /FL=cb:AF016369.1 qb:082756.1 qb:NN_004697.1
gb:NM_005969.1 /DEF-Homo sapiens nucleosome assembly protein 1-like 4 (NAPIL4), mRNA. /FEA-mRNA /GEN-NAPIL4 / PROD-nucleosome assembly protein 1-like 4 /DB_XREF-gi:5174612 /UG-Hs.78103 nucleosome assembly protein 1-like 4 / FL-gb:U77456.1 gb:NM_005969.1
ATPa
Consensus includes gb:A1911687 /FEA=EST /DB_XREF=g1:5631542 /DB_XREF=est:wc71g01.x1 /CLONE=IMAGE:2324112 /UG=Hs.30213 . ceroid-lipofuscinosis, neuronal 5 /FL=gb:AF068227.1 gb:NM_006493.1
eine methyltransferase red synthase reductase, isofoi se reductase /FL=gb:NM_024
gb:NM_006833.1 /DEF=Homo sapiens COP9 subunit 6 (MOV34 homolog, 34 kD) (MOV34-34KD), .mRNA. /FEA=mRNA /GEN=MOV34-34KD / PROD=COP9 subunit 6 (MOV34 homolog, 34 kD) /DE_XREF=gi:5803095 /UG=Hs.15591 COP9 subunit 6 (MOV34 homolog, 34 kD) / PE_ED=ED:BC002520.1 gb:U70735.1 gb:NM_006833.1
sapiens HSPC160 protein (HSPC160), mRNA. /FEA-mRNA /GEN-HSPC160 /FROD-HSPC160 protein / s.13144 HSPC160 protein /FL-gb:AF161509.1 gb:NM_014182.1
gb:NM_005770.1 /DEF=Homo sapiens small EDRK-rich factor 2 (SERF2), mRNA. /FER=mRNA (GEN=SERF2 /FRO)=small EDRK-rich factor 2 /FL=gb:AF320073.1 gb:AF073298.1 gb:NM_005770.1

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	Consensus includes gb:AV701283 /FEA-EST /DB_XREF=gi:10717613 /DB_XREF=est:AV701283 /CLONE=ADAAGD06 /UG=Hs.50785 SEC22; vesicle
209206 at	trafficking protein (S. cerevisiae)-11Ke 1 /rL=g0:bCU0120%:1 g0:nrc=r=r=r=r=r=r=r=r=r=r=r=r=r=r=r=r=r=r=
212165_at	consensus includes guistavious). List includes guistavious for the sequence sapiens clone 24606 mRNA sequence
208983 s at	gb:N37780.1 /DEF=Human leukocyte surface protein (CD31) mRNA, complete cds. / FEA-muna / VEA-D31 / NT-D31 / DEF=Human leukocyte surface protein (CD31 anison / FE-D31 / NT-D31 / DG-Hs. 78146 plateletendothelial cell adhesion molecule (CD31 anison / FE-D31 / DF-D31 / DR CM1401 / PDAN CM1401 / DNA CM1401 /
	gb:NM_002109.2 /DEF=Homo sapiens histidyl-tRNA synthetase (HARS), mRNA. /FEA=mRNA /GEN=HARS /FRUD=HISLIGYL LANA SYNTHER SYNTHEFEA: 6996013 /UG=HS.77798 histidyl-tRNA synthetase /FL=gb:NM_002109.2
	gb:NM_005827.1 /DEF-Homo sapiens UDP-galactose transporter related (UGTREL1), mRNA. /FEA-mRNA /GEN-UGTKEL1 / RD:NM_005827.1 /DEF-Homo sapiens UDP-galactose transporter related /FL-gb:D87989.1
202433_ac	gbing Object / DEF-Homo sapiens HS1 binding protein (HAXI), mRNA, /FEA-mRNA /GEN-HAXI /PROD-HS1 binding protein / gbing Object / DEF-Homo sapiens HS1 binding protein /FF-ch;NN 006118,2 db:BC005240:1 gb:U6856.1
201145_at	DB_XREF=g1:1.4%35353 / OG=Na:1.531
מירו בדדץ	gb:NN 003139.1 /DEF=Homo sapiens signal recognition particle receptor (docking protein) (SRPR), mRNA. /FEA=mRNA /GENESKEN/ PROD=signal recognition particle receptor (dockingprotein) /DB_XREF=gi:4507222 /UG=Hs.75730 signal recognition particle receptor (dockingprotein) /DB_XREF=gi:4507222 /UG=Hs.75730 signal recognition particle receptor (dockingprotein) /DB_XREF=gi:4507222 /UG=Hs.75730 signal recognition particle receptor
200918_s_at	[docking protein] /Elgasocourgass Bosne Control PPARBP JENA-MENA / GEN-PPARBP / PROD-thyroid hormone receptor gbs.NN 004774.1 / DEF-Homo sapiens PRODE PRODE Jena PROD
203497_at	interactor 2 /DB_XREF=g1:4/59265 /UG=HS.15569 Frak Dinding process /FEA=mRNA /GEN=TOR3 /DB_XREF=g1:339477 /UG=HS.101047 /UD-MS.M31523.1 /DEF=Human transcription factor (E2A) mRNA, complete cds. /FEA=mRNA /GEN=TOR3 /DB_XREF=g1:339477 /UG=HS.101047
209153_s_at	ranscription factor 3 (E2A immunoglobulin enhancer binding factors E1284) (Firegis 13.23. complex submit 5 (MGC3038), mRMA. /
•	gb:NM 030978.1 /DEF-Homo sapiens hypothetical protein similar to actin relatedprotein 23 complex, subunit 5 /DB XREF-gi:13569955 / FFA-mRNA /GEN-MGG3038 /PROD-hypothetical protein similar to actin relatedprotein 23 complex,
220966_x_at	FL-gb:NM_030978.1 Consensus includes gb:AA160181 /FEA=EST /DB_XREF=g1:1734757 /DB_XREF=est:zo81d09.s1 /CLONE=IMAGE:593297 /
212785_s_at	UG=Hs.278635 HDCMA18P protein
219176 at	
1 - 1	JOC51160 /PROD=VFSZ8 process /
2186/9_s_ac	DP_ANGE_91://JOSE=HOMO Sapiens chemokine (C-C motif) receptor 5 (CCR5), mRNA. /FEA-mRNA /GEN=CCR5 /PROD=chemokine (C-C motif) gb:NM_000579:1 /DEF=Homo sapiens chemokine (C-C motif) receptor 5 /FE=chiU54994:1 qb:U57840:1 gb:NM_000579:1
206991_s_at	il E r
209770_at	UG-HS.284283 butyrophilin, subramly 3, member Al /FL-guidous Consensus includes gb:AL042220 /FEA-EST /DB_XREF=gi:5421564 /DB_XREF=est:DKRZp434H1920_s1 /CLONE-DKFZp434H1920 /UG-Hs.141727
214268_s_at	myotubularin related protein 4
	gb:NM_000398.3 /DEF=Homo sapiens diaphorase (NADA) (Cyconinume D-) Ecutemes (NED) (Cyconinum / Cyconinum / Cyconin
201885_s_at	
209688_s_at	DB_XREF=g1:13477224 /UG=HS.26118 Homo sapiens clone 24/66 mkNA Sequence /rb=gurecoscoscoscoscoscoscoscoscoscoscoscoscosc
203487_s_at	gb:NM_015396.1 / DEF=HGMD SEQIENS DREZP434A043 protein /FL=gb:NM_015396.1 / DB_XREF=gi:7661561 / UG=Hs.102708 DREZP434A043 protein / CH_GB:NM_015396.1 / DB_XREF=gi:7661561 / UG=Hs.102708 DREZP434A043 protein / CH_GB:NM_015396.1 / CH_GB:NM_015396.
202155_s_at	gb:NM_005085.1 /DEF=Homo sapiens nucleoporin 214kD (CAIN) (NUF21#) unvan. /Linearin / / / / / / / / / / / / / / / / / / /
221786_at	Consensus includes gb:BF197222 /FEA=EST /DB_XKEK=g1:11083900 /DB_AKEK=est: Amount : 1.52629 KIAA0179 clone 24538 mENA sequence
212846_at	Consensus includes gb:AA811192 /FEA=EST /DB_XREF=g1:2880803 /Db_AkkF=est:uD/AD00:31 /CLCAL protein

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209444_at	21159.1 001417. ic trans
211937_at	factor 4B /FL=gb:NM_001417.1 gb:NE185696.1 /DEF=Homo sapiens oxysterol-binding protein 1 (OSBP1) mRNA, complete cds. /FEA-mRNA /GEN=OSBP1 / PROD=oxysterol-binding protein 1 /DB_XREF=gi:10441379 /UG=Hs.24734 oxysterol binding protein /FL=gb:AF185696.1 gb:M86917.1
	gb:NM_002556.1 gb:BC005903.1 /DEF-Homo sapiens, polymerase (RNA) II (DNA directed) polypeptide L (7.6kD), clone MGC:14494, mRNA, complete cds. / gb:BC005903.1 /DEF-Homo sapiens, polymerase (RNA) II (DNA directed) polypeptide L (7.6kD) /DB_XREF-gd:13543491 /FL=gb:BC005903.1
at	/gb=AL044097 /
05035_ac	ug-na.23500 / Inference apiens RNA, U transporter 1 (RNUT1), mRNA. /FEA-mRNA /GEN-RNUT1 /PROD-RNA, U transporter 1 / DB XREF=ri:5031832 /UGE-Hs.21577 RNA, U transporter 1 /FL-gb:AF039029.1 gb:NM_005701.1
221819_at	Consensus includes gb: BF791960 /FEA=EST /DB_XREF=gi:12097014 /DB_XREF=est:602252342F1 /CLONE=LMAGE:4344539 /UG=As:34500 Homo sapiens cDNA FLJ10447 fis, clone NT2RP1000851
203912_s_at	gb:NM_006730.1 /DEF=Homo sapiens deoxyribonuclease I-like 1 (DMASELL), mkNA. /Firstrucky ven-Lukella 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.
216996_s_at	Consensus includes gb:AK021557.1 /DEF=Homo sapiens CDNA FL011495 fis, Clone HEMBALU01399, mighing Saminarian Per
202138 x at	sapiens JTV1 gene (JTV1), mRNA. /FFA=mRNA /GEN=JTV1 /PK. b:U24169.1 gb:BC002853.1
219007 at	(FLJ13287), mRNA. /FEA=mKNA protein FLJ13287 /FL=gb:NM
	gb:NM_018398:1 /DEF=Homo sapiens calcium channel alpha2-delta3 subunit (HSAZ/12288), meun: /renamen alpha2-delta3 subunit /DB_XREF=g1:8923764 /UG=Hs.22958 calcium channel, voltage-dependent, alpha 2delta3 PROD=calcium channel alpha2-delta3 subunit /DB_XREF=g1:8923764 /UG=Hs.22958 calcium channel, voltage-dependent, alpha 2delta3
219714_s_at	Subunit / FLE-go.nm_losso.i. gb:NM_003191.1 / DEF-Homo sapiens threonyl-tRNA synthetase (TARS), mRNA. /FEA-mRNA /GEN-TARS / PROD-threonyl-tRNA. synthetase / gb:NM_003191.1 / DEF-Homo sapiens threonyl-tRNA synthetase /Filenth-BC000517.1 db:M63180.1 db:NM_003191.1
201263_at	DB XRRF=g1:450/350 /UG=HS:04131 CHR-CONT-CANA SYNCHOLOGY CONTROL N-MYZISTOYILYZNISFERSE MRNA, complete cds. /FEA=CDS / Consensus includes gp.aF020500.1 /DEF=Hono sapiens myzistoyl CoA;protein N-myzistoyltransferase 1 / PROD=myzistoyl CoA;protein N-myzistoyltransferase /DB_XREF=g1:2760893 /UG=HS.111039 N-myzistoyltransferase 1 /
201157_s_at	PI=gb:NM_021079.1 gb:AF020500.1 gb:AF043324.1
221593_s_at	90:BCUCIOS.: I LILIANDE DE CONTROL IN TOB XREF=G1:12804504 /UG=H5.164170 vascular Rab-GAPTBC-containing /FL=g0:BCULIOS).1 PROD=Similar to Libonomal December 1. JOB XREF=G1:1126534 / JOB XREF=est:1xx50h10.s1 / CLONE-IXAGE:265219 / UG=Hs.26968 Homo sapiens
214202_at	mRNA from chromosome 5q21-22, clone:357Ex
103142 5 9\$	gb:NM 003664.1 /DEF=Homo sapiens Adaptor-related protein camples 7. CERI-APBH /FROD-adaptor-related protein complex 3, beta lsubunit /DB_XERF=gi:4501974 / CERI-APBH /FROD-adaptor-related protein complex 3, beta 1 subunit /FL=gb:U91931.1 gb:U81504.1 gb:NM_003664.1 / CRC-PABH / FS172 adaptor-related protein complex 3, beta 1 subunit /FL=gb:U91931.1 gb:U81504.1 gb:NM_003664.1
222001 x at	Consensus includes gb:AI160126 /FEA=EST /DB_XREF=g1:3693506 /DB_XREF=est:qb51h08.xl /CLONE=IMAGE:17036/9 /UG=HS.£73974 KTAA0493 protein
213374 x at	Consensus includes gb:AW000964 /FEA=EST /DB_XREF=gi:5847880 /DB_XREF=est:wr9UhlU.X1 /CLUNE=LMAKE:2457011 /CC-1111 /CC-11111 /CC-111111 /CC-1111111 /CC-11111111 /CC-111111111 /CC-1111111111
201018_at	Consensus includes gb:AL079283.1 / DEF=Homo sapiens mRNA full length insert CDNA clone EUKUIRAGE 59315.7 f.m. accordance for the following
206206_at	gb:NM_005582:1 /DEF=Homo sapiens lymphocyte antigen 64 (mouse) homolog, radioprotective, 105kD /DB_XREF=gi:5031894 GEN=LY64 /PROD=lymphocyte antigen 64 (mouse) homolog, radioprotective, 105kD /FL=gb:D83597:1 gb:NM_005582.1 /UG=Hs.87205 lymphocyte antigen 64 (mouse) homolog, radioprotective, 105kD /FL=gb:D83597.1 gb:NM_005582.1

16 995000	Consensus includes gb:AL080184.1 /DEF-Homo sapiens mRNA; cDNA DKF2p4340071 (from clone DKF2p4340071). /FEA-mRNA / In xprp-ri.575761 /HG=Hs.7089 insulin induced protein 2 /FI-gb:AF125392.1
11 11 11 11 11 11 11 11 11 11 11 11 11	gb:A1418892 /FEA=EST /DB_XREF=gi:42648
212134_S_at	gene product gb:NW 07063.1 /DEF=Homo sapiens.vascular Rab-GAPTBC-containing (VRP), mRNA. /FEA=mRNA /GEN=VRP /PROD-vascular gb:NW 007063.1 /DEF=Homo sapiens.vascular Rab-GAPTBC-containing /FL=dp:AB024057.1 gb:NM_007063.1
204526_s_at	ne-binding protein (9.5kD) gi:7657485 /UG=Hs.3709 low
201568_at	protein (9.5KD) /FL=gp:BC001390.1 gp:D50309.1 gp:D50309.1 gp:Nm_11402.1 CLONE=INAGE:2055326 /UG=Hs.168075 Consensus includes gp.h.107759 /FEA=EST /DB_XREF=est:tb24g08.x1 /CLONE=INAGE:2055326 /UG=Hs.168075
221829_s_at	Karyopherin (importin) Deta 2 gb:NM_004068.1 /DEF=Homo sapiens adaptor-related protein complex 2, mu 1 subunit (APZM1), mRNA. /FEA-mRNA /GEN-APZM1 / PROD=adaptor-related protein complex 2, mu 1 subunit /DB_XREF=gi:4757993 /UG-Hs.152936 adaptor-related protein complex 2,
200613_at	12
213160_00	tional control protein 80 mRNA, complete cds. /FEA rerleukin enhancer binding factor 3, 90kD /FL=gb:A
2113/3_8_gr	o sapiens Deleted in split-handsplit-foot 1 region (DSS1), mRNA. /FEA- region /DB_XREF=gi:5453639 /UG=Hs.85215 Deleted in split-handsplit-foot
202276_ac	gg:www_00004.1 /DEF=Homo sapiens IMP (inosine monophosphate) dehydrogenase 2 (IMPDH2), mRNA. /FEA=mRNA /GEN=IMPDH2 /PROD=iMP (gb:NM_000884.1 /DEF=Homo sapiens IMP (inosine monophosphate) dehydrogenase 2 / (inosine monophosphate) dehydrogenase 2 /
201892_s_at	FL=gb:J04208.1 gb:NM_U0U884.1 gb:NW_C2043401 /DEF=Homo sapiens hypothetical protein FLJ13081 (FLJ13081), mRNA. /FEA=mRNA /GEN=FLJ13081 /FROD=hypothetical
27/302 ac	gb: BC001917.1 / DEF=Homo sapiens; malate dehydrogenase 2, NAD (mitochondrial), clone MGC:3559, mRNA, complete cds. // FEA=mRNA / PROD=malate dehydrogenase 2, NAD (mitochondrial) / DB XEEP=gi:12804928 / UG=Hs.111076 malate dehydrogenase 2,
209036_s_at	MAD
222010_at	acetyl-Coenzyme A acetyltransierase 2 (acetoacetyl Coenzyme a Linguaser) gb:NM 007371.2 (PEE-Homo sapiens bromodomain-containing 3 (BRD3), nRNA. /FEA-ERNA /GEN-BRD3 /FROD-bromodomain-containing gb:NM 007371.2 (DEE-Homo sapiens bromodomain-containing 3 /Finch:NM 007371.2 db:D26362.1
203825 at	protein 3 / Db_Aktr_gi:12400042 / DC-no.00000 promoters. protein 3 / Db_Aktr_gi:12400042 / DC-no.00000 promoters. pp.3300013 / DEF Human alpha-N-acetylgalactosaminidase mRNA, complete cds. /FER-mRNA / PROD-alpha-N-acetylgalactosaminidase was properly protein 1 gb:MG-1000011 gb:MG-10000011 gb:MG-1000011 gb:MG-10000011 gb:MG-1000011 gb:MG-10000011 gb
210555 g.at	gb:U884301. /DEF=Human transcription factor NFATx4 mRNA, complete cds. /FEA-mRNA /PROD-transcription factor NFATx4 of activated T-cells, cytoplasmic, calcineurin-dependent 3 /FL-gb:U85430.1 DB_XREF=gi:1835590 / UG-Hs.172674 nuclear factor of activated T-cells, cytoplasmic, calcineurin-dependent 3 /FL-gb:U85430.1
218370 s at	9
	DNA polymerase zeta (REV3L), a /DB_XREF=gi:4506482 /UG=Hs: gb:NM_002912.1 gb:AF179428.1
2080/0_s_ac	180 %
201272_at	gb:NM_001628.1
209824_s_at	gb:AB000812.1 /DEF-Homo sapiens mRNA for HMALLD, complete cus. /Fra-mana /Assa-mana /Assa-mana /Assa-mkrzn-SRC651 / hydrocarbon receptor nuclear translocator-like /Fi-gb:AB000812.1 gb:AF044881.1
209064_x_at	gb:AL136920.1 /DEF-Homo sapiens mRNA; CDNA DKFZp586C051 (from clone DKFZp586C031); Complete Cus. / En-mann (771-gp:AL136920.1 PROD=hypothetical protein /DB_XREF=g1:12053334 /UG-Hs.109643 polyadenylate binding protein-interacting protein / FII-gb:AL136920.1

4	1		

200759 x at	gb:NM_003204.1 /DEF=Homo sapiens nuclear factor (erythroid-derived 2)-like 1 (NPEZL1), mRNA. /FEA=mRNA /GEN=NPEZL1 / PROD= transcription factor 11 (basic leucine zippertype) /DB_XREF=gi:4505378 /UG=Hs.83469 nuclear factor (erythroid-derived 2)-like 1 / Ft=cb:NM 003204.1 ob:U08853.1
· 221787_at	Consensus includes gb: BF431618 /FEA=EST /DB_XREF=gi:11443732 /DB_XREF=est:7016e03.x1 /CLONE=IMAGE:3574349 //UG=HS.12342 Homo sapiens clone 24538 mRNA sequence
211581_x_at	gb:AF000426.1 /DEF=Homo sapiens LST1 mRNA, cLST1E splice variant, complete cds. /PEA=mRNA /GEN=LST1 /DB_XREF=gi:2145067 /UG=Hs.88411 lymphocyte antigen 117 /FL=gb:AF000426.1
37384_at	
218423_x_at	gb:NM_016516.1 /DEF=Homo sapiens tumor antigen SLP-8p (HCC8), mRNA. /FEA=mRNA /GEN=HCC8 /PROD=tumor antigen SLP-8p / DB_XREF=gi:7705396 /UG=Hs.48499 tumor antigen SLP-8p /FL=gb:AF102177.1 gb:NM_016516.1
210213_s_at	gb:AF022229.1 /DEF-Homo sapiens translation initiation factor 6 (eIF6) mRNA, complete cds. /FEA-mRNA /GEN-eIF6 / FROD-translation initiation factor 6 /DB_XREF-gi:2809382 /UG-Hs.5215 integrin beta 4 binding protein /FL-gb:AF022229.1
201598_s_at	gb:NW_001567.2 / DEF=Homo sapiens inositol polyphosphate phosphatase-like 1 (INPPL1), mRNA. /FEA=mRNA /GEN=INPPL1 / PROD=inositol polyphosphate phosphatase-like 1 /DB_XREF=gi:4755141 /UG-Hs.75339 inositol polyphosphate phosphatase-like 1 / FL=gb:NW_001567.2 gb:L24444.1
204301_at	gb:NW_014867.1 / DEF=Homo sapiens XIAA0711 gene product (KIAA0711), mRNR. / FEA=mRNR /GEN=KIAA0711 / PROD=KIAA0711 gene product / PI=gb:AB018254.1 gb:NW_014867.1
202877_s_at	Consensus includes gb:W72082 /FEA=EST /DB_XREF=gi:1382588 /DB_XREF=est:zd70c06.sl /CLONE=IMAGE:345994 /UG=Hs.97199 complement component Clq receptor /FL=gb:NM_012072.2 gb:U94333.1
208772_at	Consensus includes gb:AU160676 /FEA=EST /DB_XREF=g1:11022197 /DB_XREF=est:AU160676 /CLONE=Y79AA1002208 /UG=Hs.301226 KIAA1085 protein /PL=gb:AL136943.1
201998_at	Consensus includes gb:AI743792 /FEA=EST /DB_XREF=gi:5112080 /DB_XREF=est:wg53h11.X1 /CLONE=IMAGE:2368869 /UG=Hs.2554 sialyltransferase 1 (beta-galactoside alpha-2,6-sialytransferase) /FI=gb:NM_003032.1
208722_s_at	gb:BC001081.1 /DEF=Homo sapiens, anaphase-promoting complex subunit 5, clone MGC:2750, mRNA, complete cds. /FEA=mRNA / PROD=anaphase-promoting complex subunit 5 /DB_XREF=g1:12654502 /UG=Hs.7101 anaphase-promoting complex subunit 5 / FEA=mRNA / FEL=gb:BC001081.1 gb:BC001950.1 gb:AF191339.1 gb:NM_016237.1
55692_at	Cluster Incl. W22924:75H3 Homo sapiens cDNA /clone=(not-directional) /gb=W22924 /gi=1299757 /ug=Hs.96560 /len=792
217954_s_at	gb:NM_015153.1 /DEF=Homo sapiens XIAA0244 protein (KIAA0244), mRNA. /FEA=mRNA /GEN=KIAA0244 /PROD=KIAA0244 protein / DB_XREF=g1:7662017 /UG=Hs.78893 KIAA0244 protein /PL=gb:AF091622.1 gb:NM_015153.1
220307_at	gb:NM_016382.1 /DEF=Homb sapiens natural killer cell receptor 2B4 (CD244), mRNA. /FEA=mRNA /GEN=CD244 /PROD=natural killer cell receptor 2B4 /DB_XREF=gi:7706528 /UG=HS.157872 natural killer cell receptor 2B4 /FL=gb:AF242540.1 gb:AF105261.1 gb:AF145782.1 gb:AF107761.2 gb:AF117711.1 gb:NM_016382.1
211989_at' .	Consensus includes gb:NM_003079.1 / DEF=Homo sapiens SWISNF related, matrix associated, actin dependent regulator of chromatin, subfamily e, member 1 (SWAKCE1), mRNA. FFEA=CDS (GEN=SMAKCE1 / PROD=SWISNF related, matrix associated, actindependent regulator of chromatin, subfamily e, member 1 (DB_KREF=gi:4507088 /UG=Hs.312848 SWISNF related, matrix associated, lactindependent regulator of chromatin, subfamily e, member 1 /FEL=cpi:NM_003079.1.
. 4 508303	gb:BC005270.1 /DEF=Homo sapiens, NhDH dehydrogenase (ubiquinone) Fe-S protein 4 (18kD) (NhDH-coenzyme Q reductase), clone MGC:12313, mRNA, complete cds. /FRA-mRNA /FROD-NADH dehydrogenase (ubiquinone) Fe-S protein 4(18kD) (NhDH-coenzyme Q reductase) /DEF=gi:13528959 /UG=Hs.10758 NhDH dehydrogenase (ubiquinone) Fe-S protein 4 (18kD) (NhDH-coenzyme Q reductase) /PE-grine PF-S protein 4 (18kD) (NhDH-coenzyme Q reductase) /PE-grine PF-S protein 4 (18kD) (NhDH-coenzyme Q reductase) /PE-grine PF-S protein 4 (18kD)
201729 s at	gb:NM_014680.1 / DEF=Homo sapiens XIAA01100 gene product (KIAA0100), mRNR. /FEA=mRNR /GEN=KIAA0100 /FROD=KIAA0100 gene product / DB_XREF=gi:7661903 /UG=Hs.151761 KIAA0100 gene product /FL=gb:D470.1 gb:NM_014680.1
203530_s_at	gb:NM_004604.1 /DEP=Homo sapiens syntaxin 4A (placental) (STX4A), mRNA. /FEA=mRNA /GEN=STX4A /FROD=syntaxin 4A (placental) / DB_XREF=gi:4759185 /UG=Hs.83734 syntaxin 4A (placental) /FL=gb:BC002436.1 gb:NPC06007.1 gb:UN_004604.1
201622_at	gb:NM_014390.1 /DEF=Homo sapiens EBNA-2 co-activator (100kD) (p100), mRNA. /FEA=mRNA /GEN=p100 /PROD=EBNA-2 co-activator (100kD) /DD_XREF=gi:1657430 /UG=Rs.79093 EBNA-2 co-activator (100kD) /FL=gb:NM_014390.1 gb:UZ2055.1

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Tabelle 2: Gene aus Clusteranalyse 2

	Beschreibung der Sequenz in der GeneBank Datenbank
Affymetrix-interne	
Bezeichnung	
200654_at	
	gb:RM_014214.1 / DEF=Romo Sapreus incsitorings) - 1.0 / 1.0
203126_at	gb:NM_007150.1 /DEF=Homo sapiens zinc finger protein 185 (LIM domain) (ZNF185), mRNA. /FEA=mRNA /GEN=ZNF185 (LIM domain) /FIl=gb:NM_007150.1
203585_at	rotein 185 (LIM domain) / LB_AKkr=glicucss/1 / OG-ma: 1002 _ 1100
200	55 putative
702470_ac	
207008_at	04682 =Homo
200897_s.at	UG-HS.194431 palladin /FL-gb:AF077041.1 gb:AF151909.1 gb:NM_016081.1
205568 at	gb:NW U20980.2 / DEr=mamo sapieus aquaporin 9 /NY=97. mam: UG=HS.104624 aquaporin 9 /FL=gb:NW 020980.2 gb:AR008775.1 gb:AR016495.1
214022_s_ac	gb.ar280094.1 /DEF=Homo sapiens transcriptional coactivator Sp110b mRNA, complete cds. /FEA=mRNA gb.ar280094.1 /DEF=Homo sapiens transcriptional coactivator Sp110b mRNA, complete cds. /FEA=mRNA
209762_x_at	projectediscriptional coactivator opino per phosphorylase (UP), mRNA. /FEA-mRNA /GEN-UP /PROD-uridine phosphorylase gb:nM_003364.1 /DEF-Homo sapiens uridine phosphorylase
203234_at	/DB_XREF=g1:450/838 /UG=HS.//5/3 urigine phosphoryyanse /rings/carecarraj62b12.si /CLONE=IMAGE:1394879 Consensus includes gb:AA897514 /FEA=EST /DB_XREF=g1:3034134 /DB_XREF=est:aj62b12.si /CLONE=IMAGE:1394879
201940_at	/UG-HS.5057 carboxypeptidase D /FL=gb:U65090.1 gb:D85390.1 gb:NM_U01304.2
209310 s.at	gb:U25804.1 /DEF=Numan 1cn-2 cysteine protease arak, comprete co. 1 28978.1 gb:NM_001225.1 gb:U25804.1 gb:U28014.1 apsoptosis-related cysteine protease /FL=gb:U28976.1 gb:U28977.1 gb:U28978.1 gb:NM_001225.1 gb:U25804.1 gb:U28014.1
4. 80000	gb:NW_014821.1 /DEF=Homo sapiens KIAA0317 gene product (KIAA0317), mRNA. /FEA=mRNA /GEN=KIAA0317
בטבונס מר	gb:NV 006241.1 /DEF=Homo sapiens phosphoridylinositol transfer protein (PITPN), mRNA. /FRA-mRNA /GEN=PITPN / pp.NV 006241.1 /DEF=Homo sapiens phosphoridylinositol transfer protein /DB_XREF=gi:5453907 /UG=Hs.79709 phosphoridylinositol transfer protein /
201192_s_at	PL-gb: D30036.1 gb: M73704.1 gb: MX 006224.1
	gb:NN_004509.1 /DEF=Homo sapiens interferon-induced protein 41, 30kD (1F141), mann. /firshmann /orn-in-1, 30kD / PROD-interferon-induced protein 41, 30kD /DB_XREF=g1:4758585 /UG=Hs.241510 interferon-induced protein 41, 30kD /
208012_x_at	PL=gb:L22342.1 gb:NM 004509.1
20913/_s_ac	go.Nm. 016655. 17 DEF=Homo sapiens putative nuclear protein (LOC51307), mRNA. /FRA-mRNA /GEN=LOC51307 / DEF=Homo sapiens putative nuclear protein /FL=gb:AF251040.1 gb:NM_016605.1
218023_S_ac	Consensus includes gb:H15647 /FEA=EST /DB_XREF=g1:880467 /DB_XREF=est:ym27b09.sl /CLONE=IMAGE:49287 /
201190_s_at	UG=Hs.79709 phosphotidylinositol transfer process (Frages Joseph)

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-	Library Committee cds. /FEA-mRNA /	
	gb:U97075.1 /DEF=Homo sapiens FLICE-like inhibitory process State Communery CASP8 and PADD-like apoptosis regulator /	
210563_x_at	FL=gb:U97075.1	
10 500500	Consensus includes gb:BE788439 /rEa=EST /LD_AKKE-gl:ruczoco) /LD_AKKE-gl:ruczococococococococococococococococococo	
	ibroblast growth factor (acidic) intracellular binding process. 1.72623; e growth factor (acidic) intracellularbinding protein /DB_XREF=gi:72623; e growth factor (acidic) intracellular binding protein /FL=gb:AF11944.1 gb:AF171945.	•
202041_s_at	gb:AF010187.2 gb:NW_004214.3	•
201001_s_at	Consensus includes garbaraced from E2 variant 1 /FL=gb:U39361.1 gb:NM 003349.2 gb:BC000408.1 UG-Hs.75875 ubiquitin-conjugating enzyme E2 variant 1 /FL=gb:U39361.1 gb:NM 003349.2 gb:BC000408.1	
13075 at	Consensus includes gb:AB020677.2 /DEE=HOMO Sagreta Marks 101 Allean Consensus includes gb:AB020677.2 /DEE=HOMO Sagreta Marks 102 Allean December 102 ARRF=g1:6635136 /UG=HS.18166 KIAA0870 protein /DEAFA564C1940 /	
22.00	no sapiens DXFZF564C1940 protein (DXFZF564C1940), mrvn. / Fer-mrvn. / Cond424.1 gb:NM_014045.1 ein /DB_XREF=gi:13027587 /UG=HS.3804 DXFZF564C1940 protein /FL=gb:BC000424.1 gb:NM_014045.1	
201412_at	gb: AF131760.1	
203708 at	DB_XREF=gi:4505662 /UG=Hs.188 phosphodiesterase 45, CAUF-Specials (Managers)	•
1, 2, 200,000	gb:NM 014869.1 /DEF=Homo sapiens KIAA0763 gene product (ALAMO18306.1 gb:NM 014869.1 DB XREF=qi:7662289 /UG=Hs.4764 KIAA0763 gene product /PL=gb:AB018306.1 gb:NM 014869.1	
203207-8-45	ling protein ROC1 (ROC1) mRNA, complete cos. 11:4234917 /UG=Hs.96038 Ric (Drosophila)-like	
209882_at	FL-45:01123:-1 gb:N78165.1 gb:AF084462.1 gb:NM_006912.1 PL-45:01123:-1 gb:U718165.1 gb:AF084462.1 gb:NREF=gr / DB_XREF=est:601437912F1 /CLONE=IMAGE:3922971 /	
204276_at	UG-HS.274701 thymidine kinase 2, mitochondrial /FL=gb:NM 004614.1 gb:U77088.1 UG-HS.274701 thymidine kinase 2, mitochondrial /FL=gb:NM 004614.1 gb:U77088.1	
221984 s at:	Consensus includes gb:AL040896 /FEA=ESI /DE_ARGE-91.35031 / DE_ARGE-91.35031 / DEA = MRNA /GEN=PLJ22479 /	
	gb:NW 024900.1 /DEF=Homo sapiens hypothetical protein FLJ2419 (FLJ2419) (FLJ2419 /FL=gb:NM_024900.1	
218517_at	Consensus includes gb:BF593650 /FEA=EST /DB_XREF=g1:11685974 /DB_XREF=est:nac02d03.x1 /CLONE=IMAGE:32/5957 /	
208740_at	UG=HS.23964 sin3-associated polypeptide, 18kD /FL=gp:NM_UD3070.2 gp.0020227615F1 /CLONE=IMAGE:4332866 / nh.NM n04776.1	
221484 at	gg:	·
	gb:BC001903.1 /DEF=Homo sapiens, Similar to interleuvin 10 receptor, Peta-mRNA /PROD-Similar to interleukin 10 receptor, Deta /DB_XREF=gi:12804902 /UG=Hs.173936 interleukin 10 receptor, Peta-mRNA /PROD-Similar to interleukin 10 receptor, Deta /DB_XREF=gi:12804902 /UG=Hs.173936 interleukin 10 receptor,	
209575_at	beta /FL=gb:ECU11903.1 gb:RM_COUCECT. Deta (SF3B2), mRNA. /FEA=mRNA /GEN=SF3B2 / db:NM_006842.1 /DEF=Homo sapiens spliting factor 3b, subunit 2, 145kD /	
100000	/DB_XREF=gr:5803154 /UG=ns:10310 spiritums / PER=nRNA /	
	Consensus includes gb:BC001805.1 /DEF=Homo sapiens, clone IMARE:33430.0, march, processin FLJ13556 similar to N-myc PROD=Unknown (protein for IMAGE:3543670) /DB_XREF=g1:12804742 /UG=Hs.240615 hypothetical protein FLJ13556 similar to N-myc	
217286_s_at	downstream regulated 3 downstream regulated 3 downstream JERA=mRNA GEN=MIN51 JPROD=MIN51 protein	
207842_s_at	DB_XREF=gi:6678887 /UG-Hs.83422 MLMS1 protein /FL=gb:NM_007359.1	
	<pre>gb:NM_004481.2 /DEF=Homo sapiens unr-w-accty, a.p.m. prop-polypeptide N-acctylgalactosaminyltransferase 2 / (GalNAc-T2) (GALNT2), mRNA. /FEA=mRNA./GEN=GALNT2 /PROD-polypeptide N-acctylgalactosaminyltransferase 2 DB_XREF=gi:9945385 /UG-Hs.130181 UDP-N-acctyl-alpha-D-galactosamine:polypeptide N-acctylgalactosaminyltransferase 2</pre>	
217788_s_at	(Gainhc-T2) /FL=gb:NM_004481.2	

	11 Jense 1 Jen
48612 at	nc25b01.rl Homo sapiens cDNA /clone=IMAGE-1009129 /gp=AAZZJJJO /gl-AAZZJJJO /gl-AAZZJJJO /gl-AAZZJJJO /gl-AAZZJJJO /gl-AAZZJJJJ /gl-AAZZJJJJJ /gl-AAZZJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJ
4	Consensus includes gb: AA524505 /FEA=EST / DB_AKEF=g1:220-433 / CL_ALANGE: 3116756 / HIGHS: 321707 KIAA0742 protein
212089_s_ac	
222047_s_at	UG-Hs. 11801 argenate resistance procent and the procent and the procent activity, collagenase
	gp:NM_U0324.1 /DE-From Septem Constitute / PROD=tissue inhibitor of metalloproteinase iprecuisor / inhibitor (TIMPI), mRASA, /FRA-FRANA (CENTITAPI / PROD=tissue inhibitor of metalloproteinase (erythroid potentiating activity, collagenase
201666 at	Inhibitor) /FL=gb:BC000866.1 gb:M12670.1 gb:M59906.1 gb:NN 003254.1
	Consensus includes gb:W37431 /FER=EST /DB XREF=g1:1319023 /Jb_Ardr=cs::252211 gb:U19759.1
203218_at	UG=HS.246857 mitogen-activated process Assert Asser
213118 at	GEN-DKF27634M1526 /PROD-hypothetical protein /DB_XREF=gi:12053152 /UG=Hs.153293 KLANO/U1 protein signal), 1 (GOLGB1),
	gb:NM_004487.1 /DEF=Romo sapiens golg1 autoantigen, golg1 suntants, % margeolg1n (with transmembrane signal), 1 / men / PPA=mRNA /GEN=GOLGB1 /PROD=golg1 autoantigen, golgin subfamily b, margeolg1 (with transmembrane signal), 1 /
_	DB XREF=g1:4758453 /UG=Hs.7844 golgi autoantigen, golgin subtamily b, macrogorgin (************************************
201057_s_at	FL=gp:NM_U0446/.1 Comments includes db:BP575514 /FEA=EST /DB_XREF=gi:11649318 /DB_XREF=est:602133090F1 /CLONE=IMAGE:42880/9 /
217738 at	UG=Hs. 239138 pre-B-cell colony-enhancing factor /FL=gb:U02020.1 gb:NM_U02748.1 /CLONE=IMAGE:339167 /
	Consensus includes gb:W60953 /FEA=EST /DB_XKEK=g1:130/121 /DD_ANA 007126.2
208648_at	Consensus includes gb:AL565074 /FEA=EST /DB_XREF=gi:12916087 /DB_XREF=est:AL565074 /CLONE=CSUDNUU37840 (3 plane)
212242_at	UG-Hs.75318 tubulin, alpha 1 (testis specific)
•	Consensus includes gb:BE963444 /rEA=ESI / DE_ANALESISTED CONSENSUS includes gb:BE963444 / FEA=ESI / DE_GD:NM_020424.1
203897_ac	107116 gene, clone MGC:10940, mux, compress F=gi:13779253 /UG=Hs. 47986 Homo sapiens,
209513_s_at	2610207116
111767 c at	gb:BC005978.1 /DEF=Homo sapiens, Karyognerin argus 1, importinalpha 1) /DB_XREF=gi:13543656 /FL=gb:BC005978.1 FER=mRNA /PROD=karyopherin alpha 2 (RAG cohort 1, importinalpha 1) /DB_XREF=gi:135746651, mRNA, /FEA=mRNA /
35-0-70-117	hypothetical protein from EUKOLMA protein from EUROIMAGE 197,7056 /DE
219639_x_at	from EUROIMAGE 1977056 /FL=gb:NM_020213.1
217882 at	gb:NM U1844/11 / DEF = HGLD SEPTEM 30 kDa protein /FL=gb: AFIS7321.1 gb:NM_018447.1 (F5), mRNA. /FEA=mRNA /GEN=FS /
	ccelerin,
204714_s_at	Figb:NM_000130.2 gb:H1596.1 gb:H1595.1 gb:H1595.1 gb:BC000794.1 /DEF=H0mo sapiens, pre-mRNA, gplicing factor similar to S. cerevisiae Prp18 /DB_XREF=gi:12653992 /
10 6 7 10 10 10 10 10 10 10 10 10 10 10 10 10	complete cds. /FEA=mRNA /FROD=pre-manAs Spiriting Lactor 18 /FL=gb:BC000794.1 gb:U51990.1 gb:NA 003675.1 UG=Hs.155244 pre-mRNA processing factor 18 /FL=gb:BC000794.1 gb:U51990.1 gb:NA OFFN-MID98 /PROD=nucleoporin 98 /
- TOT77	gb:U41815.1 /DEF-Human nucleoporin 98 (NUP98) mENA, complete cds. /Fra=mkwa /can-nors / recomplete cds. / recomple
210793_s_at	DB_XREF=g1:11041/4 / OG=ns.11223
	841445.
202564 s at	ke 1 /FL-gb: BCO00642.1 gb: AF063605.1 gb: AF161461.1 gb: NM 015344. ke 1 /FL-gb: BCO00642.1 gb: AF063605.1 gb: AF161461.1 gb: NM 015344.
7 PLP 10	gb:NM_018643.1, /DEF=Homo sapiens triggering receptor expressed on myelold cells1 /DB_XREF=gi:8924261 /UG=HS.283022 triggering receptor expressed on myelold cells1 /UG=HS.28302 triggering receptor expressed on myelold receptor expressed on m
757 5 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	

	4 (ARL4), mRNA. /FEA=mRNA /GEN=ARL4 /
	mmo sapiens Aur-ribosylation factor—inc 201672 ADP-ribosylation fact factor—like 4 /DB_XREF=gi:5031602 /UG=Hs.201672 ADP-ribosylation fact
205020_s_at	FL=gb:U73960.1 gb:NM_U02-301 Consensus includes gb:AB014576.1 (DEF=Homo sapiens mRNA for KIAA0676 protein, partial cds. /FEA=mRNA /
212052_s_at	-mrna /gen-svil /
202565 s at	gb:NM_003174.2 / DEFENCED Septemb Septemb Supervillin /FL=gb:NM_003174.2 gb:AF051850.1 gb:AF051651.1 pp:AF051651.1
100000	gb. AF272036.1 / DEF=Homo sapiens Rag D mRNA, complete cds. / FeA=manA / FAAD-And D / DEF=Homo sapiens Rag D mRNA 021244.1 gb: R7272036.1 gb: BC003088.1
75.1244.S.gc	gb:NW_024293.1 /DEF-Homo sapiens hypothetical protein MGC3035 (MGC3035), mRNA. FEAS-mANA (GENERALD), PROPERTY (PROPERTY OF PROPERTY OF PROPERTY (PROPERTY OF PROPERTY OF PROPE
218037_at	25.5
214107_x_at	Consensus includes guismasogo (research production) consensus and service of the construction of the const
210184 at	gb: M81695.1 /DEF=H.sapiens leukocyte adnesion glycoprotein ploto alpha polypeptide) /FL=gb: M81695.1 gb: NM_000887.2 DB_XREF=gi: 487829 /UG=Hs.51077 integrin, dlpha X (antigen CD11C (pl50), alpha polypeptide) /FL=gb: M81695.1 gb: NM_000887.2
100000	gb:U19179.1 /DEF=Human (Hin-2) mRNA, complete cds. /FeA=mankA /GEN-nin-2 /DEF=Human (Hin-2) mRNA / Complete cds. /FeA=mankA /GEN-nin-2 / Ann vorg::5114048 /
227777777	gb:AF085357.1 /DEF=Homo sapiens flotillin mRNA, complete cds. /FDA=mRNA /FNUD=IDCLILL /DD_NAL 3
208749_X_ac	05-113-11-20-11-11-20-1-20-1-20-1-20-1-20-
212561_at	Consensus includes 92.011088.1 (DEE-Homo saplens mRNN for KIAA0516 protein, partial cds. /FEA-mRNN /GEN-KLAA0510
212470_at	PROD=KIAA0516 protein /DB_XREP=gi:3043555 /UG=Hs.129872 sperm associated antigen > /CLONE=IMAGE:2446292 /
214937 x at	Consensus includes gb:Al92481/ //EA=E31 / DB_ARAF gr.3000102 / DB_ARAF g
	gb:NM 021975.1 /DEF-Homo sapiens v-rel avian reticuloendotheliosis Vikal oncogene monarco n
	Kappa 119nt polypeptide gene dimence. The control of fouclear factor of kappa light polypeptide geneenmanger in PROD-rel avian reticulosis viral oncogene homolog A properties avian 1988 (1984) by Strengther (1986) 198 St
201783_s_at	(nuclear factor of kappa light polypeptide gene enhancer in B-cells 3 (p65)) /FL=g0:NN0213/3.1 g2
	gb:NM_021188.1 /DEF=Homo sapiens clones 2360/ and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:10863994 /UG=Hs.7137 clones 23667 and 23775 zinc finger protein /DB_XREF=gi:1086394 /UG=Hs.7137 clones 23667 and 23775 zinc finger pro
202010_s_at	FI-SD:NM_021188.1 9D:000210.1 9D:NM_00054011 /DEF-And sapidate interferon induced transmembrane protein 1 (9-27) (IFIIM1), mRNA. /FEA-mRNA /GEN=FFITM. / pp:NM_00054011 //DEF-Andread ransmembrane protein 1 (9-27) //DE_XREF-gi:4504580 //UG-HS.146360 interferon induced transmembrane
201601 x at	protein 1 (9-27) /FL=gb:BC000897.1 gb:J04164.1 gb:NM_003641.1
	gb:NM_003010.1 / DEF=Homo sapiens mitogen-activated process / / DB_XREF=g1:4506888 / / UG=HS.75217 mitogen-activated protein kinase kinase 4 / DB_XREF=g1:4506888 / UG=HS.75217 mitogen-activated protein kinase kinase 4 / DB_XREF=g1:4506888 / UG=HS.75217 mitogen-activated protein kinase kinase 4 / DB_XREF=g1:4506888 / UG=HS.75217 mitogen-activated protein kinase kinase 4 / DB_XREF=g1:4506888 / UG=HS.75217 mitogen-activated protein kinase kinase 4 / DB_XREF=g1:4506888 / UG=HS.75217 mitogen-activated protein kinase kinase 4 / DB_XREF=g1:4506888 / UG=HS.75217 mitogen-activated protein kinase kinase 4 / DB_XREF=g1:4506888 / UG=HS.75217 mitogen-activated protein kinase 4 / DB_XREF=g1:450688 / UG=HS.75217 mitogen-activated
203266_s_at	FL=gb:NM 003010.1 gb:L36870.1 gb:U1//43.1
219863_at	gb:NM_U16343.1 / DEF = FOUND SELECTION OF THE SELECTION O
212322 at	Consensus includes gb: BE999972 /FEAREST / DB_ARGE=91:10100220 / Consensus includes gb: BE9999972 /FEAREST / DB_ARGE=91:1010020 / DB_ARGE=186613 sphingosine-1-phosphate lyase 1 /FL=gb:AR4638.1
204206 at	gb:NM_020310.1 /DEF=Homo sapiens MAX binding protein /FL=gb:NM_020310.1 /DEX=Epi:9945317 /UG=Hs.25497 MAX binding protein /FL=gb:NM_020310.1 /UG=Hs.25497 MAX binding protein
203278 s at	gb.NM_016621.1 /DEF-Homo sapiens hypothetical protein (LOC21217), mrvr. / Location /FL=gb.AF208848.1 gb.NM_016621.1 prop-hypothetical protein /DB_XREF=g1:7706159 /UG=Hs.106826 RIAA1696 protein /FL=gb.AF208848.1 gb.NM_016621.1
202210-3-00	

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septor; coupled				•
gb:NW 014879.1 /DEF=Homo sapiens KIAA0001 gene product; putative G-protein-coupled receptor; g procession-coupled receptor; f procession-coupled receptor r	•	06637_at receptor; procedure saniens chitobiase, di-N-acetyl (CTBS), mRNA. /FPA-mRNA GENULLES /FRANCOLLES.	GD: IN COCK!	218924_s_at
		읾		21

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Tabelle 3: Gene aus Clusteranalyse 3

	A f femotion in	1
	interne Be-	beschretbung der Sequenz in der GeneBank Datenbank
	zeichnung	
	36711_at	Cluster Incl. AL021977:bK447C4.1 (novel MAPP (v-maf musculoaponeurotic fibrosarcoma (avian) oncogene family, protein F) E protein) /cds=(0.494, /ch=Al021977 /rl=491475 /rr=Fr 51305 /lon=110
	210845_s_at	gb:U08839.1 /DEF=Human urokinase-type plasminogen activator receptor montes complete cds. /FEA=mRNA /FROD=urokinase-type plasminogen activator receptor /UG-Hs. 17657 nls=minogen activator receptor /DB XRRF=q1:517197 /UG-Hs. 17657 nls=minogen activator models of the control of
	202643_s_at	Consensus includes gb:A1738896 /FEA=EST / DB_XREF=gi:5100877 / DB_XREF=est:w422902.x1 /CLONE=IMAGE:2391026 /UG=Hs.211600 tumor necrosis factor, alpha-induced protein 3 /FI=ch:M946F.1 Ah.WM 006790 1
	205476_at	gb:NM_004591.1 /DEF=Homo sapiens small inducible cytokine subfamily A (Cys-Cys), member 20 (SCYA20), mRNA. /FEA=mRNA / GEN=SCXA20 /PROD=small inducible cytokine subfamily A (Cys-Cys), member 20 /DE_XRRF=91:4759075 /UG=HS.75498 small inducible cytokine subfamily A (Cys-Cys), member 20 /FL=ob:U6419.1 ch:177035 1 ch.ns. 04.85015 1 ch.ns. 04.85015
		gb:NN 004951.1 /DEF=Homo sapiens Epstein-Barr virus induced gene 2 (lymphocyte-specific G protein-coupled receptor) (EBI2), nRNA. /PEA-mRNA /GEM-EBI2 /PROD-Epstein-Barr virus induced gene 2 (lymphocyte-specific G protein-coupled receptor) / DB_XREF=gi:4826705 /UG-Hs.784 Epstein-Barr virus induced gene 2 (lymphocyte-specific G protein-coupled recentor) /
	205419_at	PL=gb:L08177.1 gb:NW_004951.1 gb:L07555.1.7DE=Homo sapiens early activation antigen CD69 mRNA, complete cds. /FEA-mRNA /PROD=early activation antigen CD69 /
	205767_at	TECHNOLOGIST 1 / DEFENDED SUIT LOUR SHILLY 1-CELL SCINGLION SHILLY 1-CELL SCINGLOOP (FLEGD: L0755.1 gb:NM 001781.1 gb:NM 001432.1 / DEFENDED SEPTEMBER OF SEPTEMBERS (FROD=spliregulin precursor / DB_XREF=g1:4557566 / DG=Hs.115263 spliregulin / PL=ch:103038.1 ch:NM 001432.1
	203821_at	gb:NM_001945.1./DEF=Homo sapiens diphtheria toxin receptor (heparin-binding epidermal growth factor-like growth factor) (DTR), DB_XNEF=91.4503412 /UG=Ms.70G=61phtheria toxin receptor (heparin-bindingepidermal growth factor-like growth factor) / TO=Ms.70G=91919 (JDH), DB_XNEF=91.4503412 /UG=Ms.70G diphtheria toxin receptor (heparin-binding epidermal growth factor-like growth factor) / TD=91.30F.NM 001945.1
	211924_s_at	9D:AX029180.1 /DEF-Homo sapiens soluble urokinase plasminogen activator receptor precursor (SUPAR) mRNA, complete eds. / FEA-CDS /GEN-SUPAR / PROD-soluble urokinase plasminogen arrivator recentorment on version and activator recentorment.
	205403_at	gb:NM_004633.1 /DEF=Homo sapiens interleukin 1 receptor, type II (ILIR2), mRNA. /FEA=mRNA /GEN=ILIR2 /PROD=interleukin 1 receptor, type II /DB_XREF=gi:4758597 /UG=Hs.25333 interleukin 1 receptor, type II /DB_XREF=gi:4758597 /UG=Hs.25333 interleukin 1 receptor, type II
	204351_at	gb:NM_005980.1 /DEF=Homo sapiens S100 calcium-binding protein F (S100P), mNAS. /FEA=mRNA /GEN=S100P /PROD=S100 calcium-binding protein P /PL=ch:NM_005980.1
	206115_at	gb.NM_004430.1 /DEF=Homo sapiens early growth response 3 (EGR3), mRNA. /FEN=mRNA /GEN=EGR3 /FROD=early growth response 3 / DB_XREF=gi:4758251 /UG=Hs.74088 early growth response 3 /FL=gb:NM_004430.1
	204103_at	gb:NM_002984.1 /DEF=Homo sapiens small inducible cytokine A4 (homologous to mouse Mip-lb) (SCVA4), mRNA. /FEA-mRNA / GEN=SCYA4 /PROD=small inducible cytokine A4 (homologous to mouseMip-lb) /DE XREF=gi:4506844 /UG=Hs.75703 small inducible Cytokine A4 (homologous to mouse Mip-lb) /Fi=gb:J04130.1 gb:NM_002984.1 gb:MZ3502.1 gb:MZ5316.1
	206522_at	gb:NM_004668.1 /DEF=Homo sapiens maltase-glucoamylase (alpha-glucosidase) (MGAM), mRNA. /FEA=mRNA /GEN=MGAM / PROD=alpha-glucosidase /DB_XREF=gi:4758711 /UG=Hs.122785 maltase-glucoamylase (alpha-glucosidase) /FL=gb:AF016833.1 gb:NM_004668.1
	202147_s_at	gb:NM_001550.1 /DER=Homo sapiens interferon-related developmental regulator 1 (IFRD1), mRNA. /FER=mRNA /GEN=IFRD1 / PROD=interferon-related developmental regulator 1 /DB_XREF=gi:4504606 /UG=Hs.7879 interferon-related developmental regulator 1 /FL=gb:BC001272.1 gb:NM_001550.1
	206515_at	gb:NM_000896.1 /DEF=Homo sapiens cytochrome P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) (CYP4F3). MRNA. /FEA=MRNA /GEN=CYP4F3 /FROD=cytochrome P450, subfamily IVF, polypeptide 3 /DB_XREF=gi:4503240 /UG=Hs.106242 Cytochrome P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) /FL=gb:AB002454.1 gb:D12620.1 gb:NK_000896.1
لتنب	204614_at	gb:NM_002575.1 /DEF=Homo sapiens serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 2 (SERPINE2), mRNA / FEA-mRNA /GEN=SERPINE2 /PROD=serine (or cysteine) proteinase inhibitor, cladeB (ovalbumin), member 2 /DE_XREF=gi:4505594 /

	inhihitor clade B (ovalbumin), member 2 /FL-gb:J02685.1 gb:J03603.1
	UG=HS.75716 Serine (or cysterne) processes and compared to the compared of the compared of the compared of the compared to the
	gb:U12767.1 /DEF=Human mitogen induced nuclear orphan receptor (minos) amon, Compress receptor subfamily 4, group A, pron-mitogen induced nuclear orphan receptor./DB_XREF=gi:924281 /UG=Hs.80561 nuclear receptor subfamily 4, group A,
209959_at	member 3 /FL-gb:U12767.1 Conseques includes gpish26328.1 /DEP=Homo sapiens mRNA; cDNA DKF2p564M2422 (from clone DKF2p564M2422); partial cds. / Conseques includes gpish2630 (PROD=btvoothetical protein /DB_XRRF=g1:4914612 /UG=HS.306320 Homo sapiens mRNA; cDNA DKF2p564M2422
215078_at	FERNMAN / VENA-125. 25.01.01.01.01.01.01.01.01.01.01.01.01.01.
201890_at	consensation reductase MZ polypeptide /FL=gb:NM_001034.1 ribonesial MGC:11022, mRNA, complete cds. /FEA=mRNA / ribonesial reductase MZ polypeptidylprolyl isomerase F (cyclophilin F), clone MGC:11022, mRNA, complete cds. /FEA=mRNA / ob. EC005020.1 /DEF=Homo sapiens, peptidylprolyl isomerase F (cyclophilin F) /
201489_at	PROD-peptidylprolyl isomerase F (cyclophilin F) /DB_XREF=g1:134//120 /OG=ns.1/312 pcptidylprolyl isomerase F (cyclophilin F) /DB_XREF=g1:134//120 /OG=ns.1/312 pcptidylprolyl isomerase F (cyclophilin P) / FEA-mRNA / GEN=THED / PROD-thrombomodulin /DB_XREF=g1:4507482 /
203888_at	gb:NM 000361.1 / DEFENGED SABPENS CHICAGOST.1 GD:NM 000361.1 OGENE-RGC32 / PROD-RGC32 protein / DB_XREF-g1:7662650 / OGENE.2030 thrombomodulun samiens RGC32 protein (RGC32), mRNA. /FEA-mRNA /GEN-RGC32 / PROD-RGC32 protein / DB_XREF-g1:7662650 /
218723_s_at	go:NM_014039:1, / LatLoam
206834_at	gb.Nr. 000313.2 / Def all a PL=gb:NM 000519.2 / 103-Hs.36977 hemoglobin, delta PL=gb:NM 000519.2 / 103-Hs.36977 hemoglobin, delta PL=gb:NM 000519.2
205479_s_at	gb:NM_002658.1 /DEFenomo Septems Francisco Alexandrogen activator, urokinase /FL=gb:MI5476.1 gb:NM_002558.1 urokinase /DB_XREF=gi:4505862 /UG=HS:7174 plasminogen activator, urokinase /PROD=adrenomedullin /DB_XREF=gi:4501944 /
202912 at	gb:NM_001124.1 /DEF=Homo sapiens acremumentiii (2021)
	89
208869_s_at	cDNA DKFZp564N1272 (from clone DkFZp5e4N12/2); Compress (LIMK2), transcript variant 2a, mRNA. /FEA=mRNA /GEN=LIMK2 / db:NN 005569:2 /DEF=Homo sapiens LIM domain kinas 2 (LIMK2), transcript variant kinase 2 /FL=gb:D45906.1 gb:NM 005569.2
202193_at	PROD-LIM domain kinase 2 isoform 2a /DB_XREK-g1:8021613 /UG-ns.c./vor.c. /FEA-mRNA /DB_XRER-g1:3387938 /UG-Hs.29206
214696_at	Homo sapiens clone 24659 mRNA sequence Homo sapiens clone 24659 mRNA sequence
38037_at	Cluster Incl. M602/8: Human neparatur Distriction of the Company of the Company of Company Similar to Homo sapiens mRNA ug-Hs. 799 /len=2242
212723_at	Consensus includes gb:AKU1/BU:1 /DEF-MUM: Septembly Septembly 10431034 /UG-Hs:72660 phosphatidylserine receptor for KIAA0585 protein. /FEA-mRNA /DB_XREF-gi:10433034 /UG-Hs:72660 phosphatidylserine receptor for KIAA0585 protein. /FEA-mRNA /DB_XREF-gi:10433034 /UG-Hs:72660 phosphatidylserine receptor
	gb:NM_006061.1 /DEF=Homo sapiens specific granular process. GEN=SGP28 /FRO=specific granula protein (28, Da); cysteine-richsecretory protein-3 /DB_XREF=g1:5174674 /UG=HS.54431 Specific GEN=SGP28 /FRO=specific granula protein (28, Da); cysteine-richsecretory protein-3 /FL=gb:NM_006061.1
207802_at	tein, complete cus. hCREM 2alpha-b prot
209967_s_at	n. N. B. XREF-gi. 532036 / UG-Hs. 155924 caMP responsive element modulator.
. 210119 at	90:U/13.1. / Une
1 1000	gb:NM_007287.1 / DEF=Homo sapiens membrane metallo-endopeptidase / DB_XREF=g1:6042199 / NG=R3:1298 . LEASERS.1298 . LEASERS.12
203433 S. A.	Consensus includes gb:AII33353 /FEA=EST /DB_XREF=g1:6360669 /DB_XREF=est:HAI95 / VUE=RB:A30cot MyOs.L., -SerContaining ICE.
213513 X at	ciated Kinase mRNA, complete cus. ptor-interacting serine-threonine k
209545_s_at	gb: AF027706.1 gb: AF064824.1 gb: AF078530.1 gp: NM_U03621.1 Fonsensus includes gb: NM_005564.1 / DEF=Homo sapiens lipocalin 2 (oncogene 24p3) (LCN2), mRNA. /FEA=CDS /GEN=LCN2 /
. 212531_at	PROD=lipocalin 2 (oncogene 24p3) /DB_XREF=g1:5031852. /UG=Hs.204238 lipocalin 2 (oncogene 2-p3)

208470_s_at	gD:NM_UZU995.1 /DEF=Homo sapiens haptoglobin-related protein (HPR), mRNA. /FEA=CDS /GEN=HPR /PROD=haptoglobin-related protein / DB_XREE=g1:10337588 /UG=HS.328822 haptoglobin-related protein /FL=ch:NM 020995.1	
200733_s_at	gb:U48296.1 /DEF=Homo sapiens protein tyzosine phosphatase PTPCAAXI (hPTPCAAXI) mRNA, complete cds. /FEA-mRNA /GEN-hPTPCAXI / PROD-protein tyrosine phosphatase PTPCAAXI /DB_XREF=gi:177754 /UG=HS.227777 protein tyrosine phosphatase type IVA, member 1 / FI=gb:U48296.1 qb:NM 003463.1	
211372_s_at	gb:U64094.1 /DEF=Human soluble type II interleukin-1 receptor mRNA, complete cds. /FEA=mRNA /FROD=soluble type II interleukin-1 receptor for 17 /Fi=nh.HF4004 1	•
204794_at	gb:NM_004418.2 /DEF=Homo sapiens dual specificity phosphatase 2 (DUSP), mRNA. /FRA=mRNA /GEN=DUSP2 /PROD=dual specificity phosphatase 2 /DE_ARRES_CI_12707563 /UG=Hs_1183 dual specificity phosphatase 3 /DE_ARRES_CI_1270764 /UG=Hs_1183 dual specificity phosphatase 3 /DE_ARRES_CI_127076 /UG=Hs_1183 dual specificity	
206177_s_at	gb:NM_000045.2 /DEF=Homo sapiens arginase, liver (ARG1), mRNA. /FEA=mRNA /GEN=ARG1 /FROD=arginase, type I /DB_XREF=gi:10947138 / UG=Hs.289057 arginase, liver /FL=gb:NM_000045.2 gb:M14502.1	
205239 at	gb:NM_001657.1 /DEF=Homo sapiens amphiregulin (schwannoma-derived growth factor) (AREG), mRNA. /FEA=mRNA /GEN=AREG / FROD=mahiregulin (schwannoma-derived growth factor) /DB_XREF=gi:4502198 /UG=Hs.270833 amphiregulin (schwannoma-derived growth	
	s gring_coross.in no sapiens interleukin 8 receptor, alpha 500 /UG=Hos 194778 interleukin 8 receptor	
200776_s_at	Consensus includes gb:AL518328 /FEA=EST /DB_XREF=est:AL518328 /CLONE=CS0DA009XR18 (3 prime) /UG=HS.155291 KIAA0005 gene product /FL=cb:D13630.1 ch:NM 0.14670 1	
215009_s_at	Consensus includes gb:U92014.1 / DEF-Human clone 121711 defective mariner transposon Hsmar2 mRNA sequence. / FEA-mRNA / DB_XREF-g1:2052456 /UG-Hs.153527 Homo sapiens uTMS:mariner-like transposon mann partial sequence. / FEA-mRNA /	·
211302_s_at	gb:L20966.1 / DEF-Human phosphodiesterase mRNA, complete cds. /FEA-mRNA /FROD=phosphosphodiesterase / DE_XREF=gi:347121 / UG=Hs.188 phosphodiesterase 48, cAMP-specific (dunce (Drosophila)-homolog phosphodiesterase 84), rech.1.3066.1	
210512_s_at	gb:AF022375.1 /DEE-Homo sapiens vascular endothelial growth factor mRNA, complete cds. /FEA-mRNA /FROD=vascular endothelial growth factor /DE_XREF=gi:3719220 /UG=HE.73793 vascular endothelial growth factor /FL=gb:M32977.1 gb:AF021375.1 gb:AF091352.1	•
212577_at	Consensus.includes gb: AA868754 /FEA=EST /DB_XREF=gi:2964199 /DB_XREF=est:ak52e09.s1 /CLONE=IMAGE:1409608 /UG=Hs.8118 KIAA0650 protein	
214866_at	Consensus includes gb:X74039.1 /DEF=H.sapiens mRNA for urokinase plasminogen activator receptor. /FEA=mRNA /FROD=urokinase plasminogen activator receptor /DB_XREF=g1:456192 /UG=Hs.179657 plasminogen activator irrivinses receptor.	
219228_at	gb:NM_018555.2 / DEF=Homo sapiens C2HZ-like zinc finger protein (ZNF361), mRNs. /FEA-mRNS /GEN=ZNF361 / PROD=C2HZ-like zinc finger protein / DB_XREF=gi:10092612 / UG=HS:147644 zinc finger protein 331 / FEL=dp:AF251515.2 dp:NM 018455 2 dp:AF27914R)	
201574_at	gb:NM_004730.1 /DEF=Homo sapiens eukaryotic translation termination factor 1 (ETF1), mRNA. /FEA-mRNA /GEN=ETF1 / PROD=eukaryotic translation termination factor 1 /DB_XREF=g1:4759033 /UG=Hs.77324 eukaryotic translation termination factor 1 / FL=gb:U90176.1 gb:M75715.1 db:NM_004730.1	
209498_at	Consensus includes gb:XI6354.1 /DEP=Human mRNA for transmembrane carcinoembryonic antigen BGPa (formerly TM1-CEA). / FEA=mRNA /PROD=TM1-CEA preprotein /DB_XREF=g1:37197 /UG=Hs.50964 carcinoembryonic antigen-related cell adhesion molecule 1 (biliary glycoprótein) /FL=qb:303858.1	•
207630_s_at	gb:NM_001881.1 / DEF-Homo sapiens cAMP responsive element modulator (CREM), mRNA. /FEA-mRNA /GEN-CREM /PROD-cAMP responsive element modulator / DB_XREF-g1:4503038 /UG-HS.155924 cAMP responsive element modulator / Frienh.NM n01881 1 . Ah. c.6271 1	•
210873_x_at	gb:U03891.2 /DEF-Homo sapiens phorbolin I mRNA, complete cds. /FEA-mRNA /PROD-phorbolin I /DB_XREF=gi:4895107 /UG-Hs.226307 phorbolin (similar to apolipoprotein B mRNA editing protein) /FE=ch:U03891.2	٠
204419_x_at	gb:NM_000184.1 /DEF=Homo sapiens hemoglobin, gamma G (HBG2), mRNA. /FEA-mRNA /GEN=HBG2 /PROD=hemoglobin, gamma G / DB_XREF=g1:6715606 /UG=HS.283108 hemoglobin, gamma G /FL=cp:NM 000184.1	
202988_s_at	gb:NM_002922.1 /DEF=Homo sapiens regulator of G-protein signalling 1 (RGS1), mRNA. (FEA-mRNA /GEN-RGS1 /PROD-regulator of G-protein signalling 1 /DE XREF=gi:4506514 /UG=HS.75256 regulator of G-protein signalling 1 /FI=ch:NM 002922.1	
209273_s_at	Consensus includes gb:BG387555 /FEA=EST /DB_XREF=gi:13281001 /DB_XREF=est:602412371F1 /CLONE=IMAGE:4521017 /UG=Hs.17776 hypothetical protein MGC4276 similar to CG8198 /FL=gb:AF284752.1 qb:BC002675.1	
202693_s_at	Consensus includes gb:AN194730 /FFR=EST /DB_XREF=g1:6473630 /DB_XREF=est:xn43d11.x1 /CLONE=IMAGE:2696469 /UG=Hs.9075 serinethreonine kinase 17a (apoptosis-inducing) /FL=gb:AB011420.1 gb:NM_004760.1	
205270_s_at	gb:NM_005565.2 /DEF=Homo sapiens lymphocyte cytosolic protein 2 (SH2 domain-containing leukocyte protein of 76kb) (LCP2), mRNA. /FEA=mRNA /GEN=LCP2 /PROD=lymphocyte cytosolic protein 2 /DB_XREF=gi:7382491 /UG=Hs.2488 lymphocyte cytosolic protein 2 (SH2 domain-containing leukocyte protein of 76kb) /FL=gb:NM_005565.2 gb:U20158.1	

	. PEA-mRNA /GEN-CDRNIA / CDRNIA), MRNA. /PEA-MRNA /GEN-CDRNIA /
•	gb:NM_000389.1 / DEF=Homo Saprens Cyclin Cycle (1917) / DB_XREF=gi:11386202 / UG=HS.179665 cyclin-dependent Kinase innibitor APP (1921-Cipi) / DB_XREF=gi:11386202 / UG=HS.179665 cyclin-dependent Kinase innibitor (1921-Cipi) / UG=HS.179665 cyclin-dependent Kinase innibitor (1921-Cipi) / UG=HS.179665 cyclin-dependent Kinase innibitor (1921-Cipi) / UG=HS.17966 cyclin-dependent Kinase innibitor (1921-Cipi) / UG=HS.17966 cyclin-dependent Kinase innibitor (1921-Cipi) / UG=HS.17966 cyclin-dependent Kinase innibitor (1921-Cipi) / UG=HS.1796 cyclin-dependent Kinase innibitor (1921
202284_s_at	(p21, Cip1) /FL=gb:NM_000389.1 gb:ECU01273.1 gb:ECU01273.1 gr:ECT01562 (FL721562), mRNA. /FEA=mRNA /GEN=FL721562 /FROD=hypothetical ch:NM 025113.1 /DEE=Homo sapiens hypothetical protein FL721562 (FL721562), mRNA. /FEA=mRNA /GEN=FL721562 /FROD=hypothetical ch:NM 025113.1 /DEE=Homo sapiens hypothetical protein FL721562 /FL721563 /FL7215
219471_at	protein FL721562 /DB XREF-gi:13376686 /UG-HS.288708 hypothetical protein rivation /LE-grimmal transducer CD24 / gb:MS8664.1 /DEF-Homo sapiens CD24 signal transducer mRNA, complete cds. /FBA-mRNA /FROD-signal transducer CD24 ignal transducer mRNA, complete cds. /FBA-mRNA /FROD-signal transducer CD24 in the complete cds. /FBA-mRNA /FROD-signal transducer CD24 antigen (small cell lung carcinoma cluster 4 antigen) /FL-gb:MS8664.1 gb:L33930.1
208651_x_at	gb:NW_01330.1 gb:NW_01330.1 gb:NW_01330.1
211434_s_at	receptor / DB_XREF=gi:3550066 /UG=Hs.302043 chemokine (C-C motiit) 1877639 /FPR=EST /DB_XREF=gi:5511255 /DB_XREF=est:wk95g09.x1 /CLO
204285_s_at	Consensus includes 92.32.00.13.20.13.00.10.10.10.10.10.10.10.10.10.10.10.10.
221824_s_at	Consensus includes go: Arthory (includes go: Arthory Consensus Inchides go: Consensus go: Con
219081_at	gb: NM_024668.1 / DEF=Homo sapiens hypothetical protein FL720288 /FL=gb: BC004457.1 gb: NM_024668.1
220528_at	gb: NM_018399.1 / DEF=Homo sapiens VAN3 protein (HisAz3090c), mann: / Linearing for the control of the control
	gb:NM_003059.1 /DEF=Homo sapiens solute carrier ramily 24 (Organic cationtransporter), member 4 /DB_XREF=gi:4507002 / FEA=mRNA (GEN=SIC22A4 /PROD=solute carrier family 22 (Organic cationtransporter), member 4 /FL=qb:AB007448.1 gb:NM_003059.1
205896_at	UG-HS.77239 SOLUTE CARTIEL FAMILY & (0.99mil Camplete cds. /FEA-mRNA /GEN-FKSG17 /PROD-FKSG17 /DB_KREF-g1:12276119 /
211445_x_at	UG-HS.307057 Homo sapiens FKSG17 mRNA, complete cds /ru-go:Arsissar UG-HS.307057 Homo sapiens FKSG17 mRNA /GEN-PWAIP1 /
	gb:NM_021127.1 / DEF=Homo Sapiens photoby 1.2-myristede-induced proteini / DB_XREF=gi:10863922 / UG=HS.96 phorbol-12-myristate-13-acetate-induced proteini / UG=HS.96 phorbol-12-myristate-13-acet
204286_s_at	process 1 / First Strangers First Strangers First Strangers (First Strangers
202503_s_at	DB_XKEF=G1: 001903 /00-marging_nine_F1B892 FR02399 mRNA, complete cds. /FEA-mRNA /PROD-PR02399 /DB_XKEF=G1:11433529 / G1:AF130113.
211560_s_at	US-BB: 77103 CYCOLING Sapiens, Similar to glyČogen synthase kinase 3 beta, clone MGC:1736, mRNA, Complete Cus. / gb;BC:0002511. (DEF-Homo sapiens, Similar to glyČogen synthase 3 beta /DB_XREF=g1:12652980 /UG-Hs.78802 glycogen synthase kinase 3 beta /DB_XREF=g1:12652980 /UG-Hs.78802 glycogen synthase 4 beta /DB_XREF=g1:12652980 /UG-Hs.78802 glycogen synthase 5 beta /DB_XREF=g1:12652980 /UG-Hs.78802 glycogen synthase 5 beta /DB_XREF=g1:12652980 /UG-Hs.78802 glycogen synthase 6 beta /DB_XREF=g1:12652980 /UG-Hs.78802 glycogen synthase 6 beta /DB_XREF=g1:12652980 /UG-H
209945_s_at	Frankon (1900) 1
217997_at	homology-like domain, family A, member 1 /FL=gb:NM 007350.1
	gb.BC005008.1 / DEF=Homo saplens, carcinoembryonic antigen-related cell adhesionmolecule 6 elone MGC:10467, mRNA. complete cds. /FEB=mRNA./ PROD=carcinoembryonic antigen-related cell from-specific cross reacting antigen /DB_XREF=gi:13477106 /UG=Hs.73848 carcinoembryonic antigen-related cell from-specific cross reacting antigen) /DB_XREF=gi:13477106 /UG=Hs.73848 carcinoembryonic antigen-related cell
203757_s_at	adhesion molecule 6 (non-specific cross reacting antigen) /FL=g0:BCU03008.1 g0:nicity = 3
205269_at	Consensus includes go:Allassal (February 2012) and Consensus includes go:Allassal go:Uclassal (VIG-HS: 2488 lymphocyte cytosolic protein 2 (SH2 domain-containing leukocyte protein of 76kD) /FEA-ERNA /GEN-FLJ11264
219049_at	gb:NM_018371.1 / DEF-Homo sapiens Appointation process 1200 hypothetical protein FLJ11264 /FL-gb:NM_018371.1 / PROD-hypothetical protein FLJ11264 /DB XREF-gi:348911
209396_s_at	66921
202637_s_at	Consensus includes gb:Ato08/22 /FEA=ES1 /DE_ANGE = 100000000000000000000000000000000000
205557_at	gb:NM_001725.1 / DEFENDED SADIEMS MACCELLILINGS FOR SADIEMS MACCELLS OF The CONTROL OF THE CONTR
207072 at	

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	Consensus includes gb: BE550486 /FEA=EST /DB_XREF=g1:9792178 /DB_XREF=est:7a27c01.xl /CLONE=IMAGE:3219336 / Consensus includes gb: BE550486 / FEA=EST /DB_XREF=g1:0.00000000000000000000000000000000000
202498_s_at	/UG-Hs.7594 solute carrier ramily 2 (inclinated girose), mRNA, /FER-mRNA /GEN-LOC51099 /PROD=CGI-58 protein
218739_at	/DB_XREF=91:7705770 /UG=Hs.19385 CGI_58 protein /FL=gb:NR-151816.1 gb:NR-1006.1 /DB_XREF=91:7705770 /UG=Hs.19385 CGI_58 protein /FL=gb:NR-1718 fis. clone HEMBA1005252, highly similar
212722_S_at	Consensus includes gD:AAAZI/00.1 /LEE-LACAD STEEF-GI:10433034 /UG-HS.72660 phosphatidylserine receptor Consensus mRNA for KIAA0585 protein. PEA-mRNA /DB_XREF-GI:10433034 /UG-HS.72660 phosphatidylserine receptor Consensus mRNA for KIAA0585 protein. STEEF-MRNA /DB XREF-GI:10433034 /UG-HS.72660 phosphatidylserine receptor Consensus mRNA for KIAA0585 protein.
. 40 0000	Consensus includes gb:AW576457 /FEA=EST /DB_AKKEF=g1:124/350 /DB_AKMEF=G1:12450 /DB_AKKEF=G1:124568 /UG=HS:20952 Homo sapiens clone 24411 mRWA seguence
212930_ar	db:NM. 000611.1 /DEF-Homo sapiens CD59 antigen p18-20 (antigen identified by monoclonal antibodies 10.36), but,
	E330, EL32 and G344) (CD59), mRNA. /FEA=mRNA /GEN=CD59 / FKUJ=LD53 and Lysel prof. (CD59), mRNA. /FEA=mRNA /GEN=CD59 / PKUJ=LD55 (CD59), mRNA. /FEA=mRNA /GEN=CD59 / PENSE=gi:10835164 //GE=s119663 CD59 antigen pl8-20
200985_s_at	(antigen identified by monoclonal antibodies 16.3AS, EJ16, EJ30, EL32 and GJ44) /FL=GDINE_U00011.1 95:10-10-10-10-10-10-10-10-10-10-10-10-10-1
	ğ
206026_s_at	/UG-HS.29352 tumor necrosis factor, alpha-induced protein 6 /ringse mRNA, complete cds. /FEA-mRNA /PROD-inositol
210740_s_at	gb:AF7733/2.1 / DEE=Homo Septems included in 1.2006345 / UC=Hs.6453 incsitcl 1.3,4-triphosphate 56 kinase / PL=gb:AFF.737.4-trisphosphate 56-kinase / DB_XREF=91:12006345 / UC=Hs.6453 incsitcl 1.3,4-trisphosphate 56-kinase / DB_XREF=91:12006345 / UC=Hs.612 / OC=Hs.811
202313 g at	Consensus includes gb:AA877765 /FEA=EST / UB_AKKE=g1:2300/30 / LB_AKKE Consensus includes gb:AA877765 /FEA=EST / UB_AKKE=g1:2300/30 / LB_AKKE
20,000	mRNA for Id-2H, complete cds. /FEA-mRNA /GEN-Id-2H /FRUD-1d-2H /DD_ANG-
201566_x_at	inhibitor of DNA binding 4, dominant megative for a for /clone=IMAGE-1712816 /clone_end=3 /gb=A1129310 /gi=3597824 /
44790 s at	ug=Hs.234923 /len=811
713036 c pt	Consensus includes gb:AW052084 /FEA=EST /DB_AREK=gl:3314443 /DB_AREK=gl:3314445 /DB_AR
	gb:NM_005204.1 /DEF=Homo sapiens mitogen-activated protein kinase kinase o lumboo, mann. () most kinase ki
16 2 7 CD200	FROM
22-2-22000	Gb:NM_002424.1 /DEF=Homo sapiens matrix metalloproteinlasse 8 (neultrophil collagelasse) /marcy/
. 40 00000	/FEA-mRNA /GEN-MARB / PROU-markix metalloproteinase 8 (neutrophil collagenase) /FL-gb: 005556.1 gb:NM 002424.1
25-020-02	Consensus includes gb: AA576961 /PEA-EST (DB XERF=gi:254435 / DB XKER=est:mmszdvo.si / closed-street / closed-
217996_at	pleckstrin homology-like domain, family A, member 1 (firstring) / https://doi.org/10.1016
208632_at	UG-Hs.5094 ring finger protein 10 /FL-gb:AB027196.1
	gb: NN 002935.1 /DEF-Homo sapiens ribonuclease, RNase A family, 3 (ecsinophilcationic protein) /DB_XREF-gi:4506550 / FEA-mRNA /GEN-RNASE3 /FROD-ribonuclease, RNase A family, 3 (ecsinophilcationic protein) /DB_XREF-gi:4506550 /
206851_at	UG-HS. 73839 ribonuclease, RNase A family, 3 (eosinophil catiour process) (CLONE-IMAGE:2136932 /UG-HS.1298 membrane
203434_s_at	Consensus includes grands and operations, enkephalinase, CALLA, CD10) /FL-gb:J03779.1 gb:NM U0.287.1 gb:NM - CPEA-mRNA / metallo-endopeptidase (neutral endopeptidase, repa-mRNA / metallo-endopeptidase);
	Consensus includes gb:AL110298.1 / DEr=Homo septens immas, Consensus includes gb:AL110298.1 / DESTRICTED Graces of Consensus includes gb:AL110298.1 / DESTRICTED From Front Fr
216236_s_at	transporter), member 3
	gb:NN_006123.1 DEF=Homo sapiens iduronate 2-suitatase (number 37mm 172458 iduronate 2-sulfatase (Hunter Syndrome) Company 2
206342_x_at	FROD-LAUSS6.1 gb:NM_006123.1 Fr_gb:L40586.1 gb:NM_006123.1
,	gb:NN_005239.1 /DEF-Homo sapiens V-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 /UG-HS.85146 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 v-ets avian erythroblastosis virus E26 norgenehomolog 2 /DB_XREF-gi:4885220 v-ets avian erythroblastosis virus erythroblastosis virus erythroblastosis virus
201329_s_at -	E26 oncogene homolog 2 /FL=gb:J041U2.1 gD:NR_UU32233.1
200731_s_at	Consensus includes grammary 1/FL=gp:U48296.1 gb:NM_003463.1 clone Y79AA1001402, weakly similar to Homo sapiens
	Consensus includes gb: AKO24029.1 / DEK=Homo Sapiens Consensus includes gb: AKO240297 / UG=Hs. 24719 modulator of apoptosis 1 /
212508_at	

	22 - 1 1 22 1 2 1 2 2 2 2 2 2 2 2 2 2 2
	FL=9D:AFJ03-2011 500000
208868_s_at	Consensus includes gp:sriziou (inchesione DKFZp564N1272); complete cds /Fi-gb:ALilab/No.1 gp:Arosova;: mRNA, sapiens mRNA; cDNA DKFZp564N1272 (from clone DKFZp564N1272); mRNA. sapiens mRNA; cDNA DKFZp564N1272 (from clone DKFZp564N1272); mRNA.
	gb.NM_000314.1 / DEF-Homo sapiens phosphares are the part of the p
204054_at	/UGERS.1011
212262_at	Consensus includes go: Artalogo (17 Lege) API42419.1 gb: AFI42422.1 mouse quaking QXI (KR domains rem. April 2014
208785 s at	Consensus includes gb: BE833893 (FEA-ES) (Market 1 Market 1 gb: NM 022818.2 gb: AF18341/1.1 (PYSF), mRNA. /FEA-mRNA / microtubule-associated proteins 1alB light chair 3 /FL=gb: Argirophy 28 (autosomal recessive) (DYSF), mRNA. /FEA-mRNA / microtubule-associated proteins 1alB light chair and argirophy 28 (autosomal recessive) (DYSF), mRNA. /FEA-mRNA / microtubule-associated proteins 1alB light chair and argirophy 28 (autosomal recessive) (DYSF), mRNA. /FEA-mRNA / microtubule-associated proteins 1alB light chair and argirophy 28 (autosomal recessive) (DYSF), mRNA. /FEA-mRNA / microtubule-associated proteins 1alB light chair and argirophy 28 (autosomal recessive) (DYSF), mRNA. / FEA-mRNA / microtubule-associated proteins 1alB light chair and argirophy 28 (autosomal recessive) (DYSF), mRNA. / FEA-mRNA / microtubule-associated proteins 1alB light chair and argirophy 28 (autosomal recessive) (DYSF) (DYSF) (MYSF)
	gb:NM_003494.1 /DEF=Homo sapiens dysferin, 11mD gittle muscular dystrophy 2B (autosomen recessive) (GNN=DYSF /PROD=dysferlin /DB_XREF=gi:4503430 /UG=Hs.143897 dysferlin, limb girdle muscular dystrophy 2B (autosomen recessive)
218660_at	FL=gb:aF075575.1 gb:NM 003494.1 FL=gb:aF075575.1 gb:NM 003494.1 FL=gb:aF075575.1 gb:NM 003494.1
221920_s_at	Consensus includes grant Solute carrier VIG=Hs.300496 mitochone Solute carrier VIG=Hs.300496 mitochone Solute carrier VIG=Hs.300496 mitochone VIG=Hs.300496 mito
211982_x_at	Consensus includes guraniscos (1200 Mines) protein PL123476 (PL123476), mRNA. /FEA=mRNA /GEN=FL123476
218647 g at	gb:NM_024640.1 /DEF=Homo sapiens Nypolnetron professional hypothetical protein FLJ23476 /FL=gD:NM_02450.1 PROD-hypothetical protein FLJ23476 /DB_XREF=gi:13375875 /UG=Hs.46736 hypothetical protein FLJ23476 /DB_XREF=gi:13375875 /UG=Hs.46736 kinase kinase kinase 4 (MAP4K4), mRNA.
	gb:NW_004834.1 /DEF=Homo sapiens mitogen-activated protein kinase kinasekinase 4 /DB_XREF=g1:4758523 PROD=mitogen-activated protein kinase kinase kinasekinase 4 /DB_XREF=g1:4758523
20657i_s_at	//CG=HS.3628 mitogen-activated protein kinase kinase kinase kinase 4 / FL=gD:AKONON: /FEA=mRNA /GEN=HBEX2 / HBEX2 / MRNA. /FEA=mRNA /GEN=HBEX2 / HBEX2 /
	gb:NM_018476.1 / DEF=Homo sapiens undergrand the control of the co
218332_at	FI-3D: AF220189.1 gb:NM_018476.1 gD:AF183410.1 gb:AF183410.1 gb:AF183410.1 member 2) (TNF), mRNA. /FEA-mRNA /GEN=INF / LAND AF1870 AND AF1870 A
	PRODetumor necrosis factor (cachectin) /DB_XREF=g1:10835154 /UG-ms.zaro/c
207113_s_at	FL-EGINAL VOOD-37-1. FL-EGINAL VOOD-37-1. FL-EGINAL VOOD-37-1. IDEPENDENS, SIMILAR to POLY (A) POLYMERERS ALPHA (FL-EGING) TOLY (A) POLYMERERS ALPHA (FL-EGING) I.
209388_at	PERA-mRNA /PROD-Similar to poly (A) polymerase /DB XREF=g1:1.1034110 /UG-18:1000 F-03111 Sequence 2 (EMR2), mRNA.
	gb:NX_013447.1 / DEF=Homo Saprems eg
207610_s_at	/UG=Hs.137354 egf-like module concaining, machine binding protein 1 (CHD1), mRNA. /FEA=mRNA /GEW=CAD1.
0 U	PROD-chromodomain helicase DNA binding protein 1 / DB XKEF=glit357445 / OF PER MIRZ hinding protein 1 /FI=gb:AF006513.1 gb:NM_001270.1
18 CONTO	gb:NN_005439.1 /DEF=Homo sapiens myeloid leukemia factor / (mis.) intermit factor / (DEF=Homo sapiens myeloid leukemia factor / DEF=Homo sapiens myeloid leukemia factor / DEF + February / DEF +
200948_at	/FL=gb:BC000898.1 gb:BC002340.1 gb:U57342.1 gb:Ar0/0535.1 gb:Nr Complete cds. /FEA=mRNA
211506_s_at	gb:AF043337.1 (DEE=Hound September Anticare Anti
217966 s.at	gb:NN 022083.1 / DEF=Homo sapiens niban protein (NIESD)
214151 s at	Consensus includes gb:AU144243 /FEATES1 /DD_AUL. 9. //UG=Hs.247118 phosphatidylinositol glycan, class aroduct of calactocerebrosidase, complete cds.
	9b:D25284.1 /DEF-Human mRNA for alternative spiled produced for the production of the properties of th
211810_s_at	product of galactocerebrosidase /DB_XREF=g1:45/445 /UG-BB:275 (MMP25)
207890 c at	SUMMANDS /PROD-matrix metalloproteinase 25 preproprotein /DB_Andr-y-1-100.
401020 B.	

	Consensus includes gb:AU145019 /PEA=EST /DB_XREF=gi:11006540 /DB_XREF=est:AU145019 /CLONE=HEMBA1003646
213056_at	/UG-HS.96427 KIAA1013 protein
10000	Consensus includes gb:AKU24214.1 / DEF-mum Saprems Com: 125.18827 KIAA0849 protein
78-78777	3E966299
214784 x at	/UG=Hs.70500 KIAA0370 protein /ros verye-e4.5811756 /DB XRRF=est:wr83e06.xl /CLONE=IMAGE:2494306
222035 S AF	Consensus includes go: Al2044/7 / Ltd-Es1 / DE_Attil -gi: Oct. Consensus includes go: Al204/8 Consensus alpha
20-0-0-0-0	Consensus includes gb:D38521.1 /DEF=Human mRNA for KIAA0077 gene, partial cds. /FEA=mRNA /GEN=KIAA0077
212219_at	/DB_XREF=gi:559329 /UG=HS.112330 AIANUV/ PLOCETH Concerns includes ob: AU144243 /FEA=EST /DB_XREF=gi:11005764 /DB_XREF=est:AU144243
214152_at	/CLONE-HEMBA1001328 /UG=Hs.247118 phosphatidylinositol glycan, class B
1 870000	gb:NM_000877.1 /DEF=Homo sapiens interleukin 1 receptor, type 1 /FL=gb:MZ7492.1 gb:NM_000877.1 receptor, type I /DB XREF=qi:4504658 /UQ=Hs.82112 interleukin 1 receptor, type I /FL=gb:MZ7492.1 gb:NM_000877.1
25-03-03-03-03-03-03-03-03-03-03-03-03-03-	ar.
207978_s_at	group A, member 3 (FL=gb:NM, 006911. gb:n/182/91. group A member 3 (FL=gb:NM, complete cds.
	gb:AF087853.1 /DEF=Homo saptens growth arrest and DNA damage inducible proteinbeta /DB_XREF=gi:12061050 /FEA=MRNA /GEN=GADD45B /PROD=growth arrest and DNA damage inducible proteinbeta /DB_XREF=gi:12061050
209304_x_at	/UG=Hs.1105/1 growth arrest and bar damage interpreted from the following includes gb:BG177920 /FEA=EST /DB_XREF=g1:12684623 /DB_XREF=est:602327839F1 /CCONE=IMAGE:4429256
214783_s_at	/UG=HS.75510 annexin All
45 05034	Cluster Incl. AL039447.DKFZp434N1010_s1 Homo sapiens.cDNA, 3 Hud /crome-DALSpronted / AL039447.DKFZp4374N1010_s1 Homo sapiens.cDNA, 3 Hud /crome-DALSpronted / AL039447.DKFZp4371DFZP4371DFZP4371DFZP4371DFZP4371DFZP471DFZP471DFZP471DFZP471DFZP471DFZP471DFZP471DFZP471DFZP471DFZP471DFZP471
40210_ar	Consensus includes gb: BG287153 /FEA=EST /DB_XREF=g1:13040709 /DB_XREF=est:602381868F1 /CLONE=1MAGE:443533
221760_at	/UG=HS.25251 mannostudse, digus, class 17, mannostudes cds. /FEA=mRNA /GEN=ILIA /PROD=interleukin 1-alpha db:MI5329.1 /DEF=Human interleukin 1-alpha (ILIA) mRNA, complete cds. /FEA=mRNA /GEN=ILIA /PROD=interleukin 1-alpha
210118_s_at	/DB_XREF=g1:186277 /UG-HS:1722 interleukin 1, alpha /Figerpointisserseriemity member 6 (TMFRSF6), mRNA. /FEA-mRNA
	gb:NM_000043.1 / DEF-Homo sapiens tumor necrosis factor receptor superfamily, /GEN-TWIRSF6 /PROD-apoptosis (APO-1) antigen 1 / DB_XREF-gi:4507582 /UG-HS.82359 tumor necrosis factor receptor superfamily,
204781_s_at	member 6 (FE-gb:N6/454.1 gb:NM_U0Uv45.1 gb:NM_UUV45.1 deltor 4 (40kD) (NCF4), transcript variant 2, mNNA.
1	gb:RM_013410.1 Ubstandin Saptems : court. of the court
207677_s_at	/ucamil.120322 metricognist arecognistic functions and dipplosphate linked moiety X) -type moiif 4 (NUDA)4, mRNA 4, mRNA 4 (NUDA)4 mRNA 4 (NUDA)4 mRNA 4 (NUDA)4 mRNA 4 (NUDA)4 mRNA 4 (NUCAME 9218)
	FER-mRNA /GEN-NUDTA / PROD-nudix (nucleoside diphosphate linked moietyx) -type motif 4 /DE XKEF = 91:10001337 DG-13:13153.1
206302_s_at	nuclised of the control of the contr
219460_s_at	protein PLJ20507 /DB_XREF=gi:8923465 /UG=Hs.202955 Aypochetical protein PLJ20507 /LEGNE=CSDA007YB20 (5 prime)
212225_at	Consensus includes garanged initiation factor (VG=Hs.150580 putative translation initiation factor factor)
204479 at	9b:NN 012383.1 / DBF-Homo sapiens osteoclast stimilating factor 1 / FL-qb:U63717.1 gb:NN 012383.1 stimilating factor 1 / DB_XREF-gi:6912563 / UG-Hs.95821 osteoclast stimilating factor 1 / PDB_XREF-gi:6912563 / UG-H
	Consensus includes gb:AK026080.1 /DEF-Homo sapiens cDNA: FLJ22427 fis, clone AKC09013. /FEA-MANA /22_ALL - 308883 enjejne factor 3a, subunit 1, 120kD
21645/_S_at	Consensus includes gb:AM293356 /PEA=EST /DB_XREF=gi:6699992 /DB_XREF=est:UI-H-BI2-ahl-c-11-0-UI.s1 /CLONE=IMAGE:2/2/20 //20 //20 //20 //20 //20 //20 //
212676_at	UG-HS.58220 Homo sapiens cDNA: FLJ23005 fis, clone LNGU0396, nighty binited to receive 1 / Nnep-Homo saniens nucleobindin 2 (NUCB2), mRNA: /FEA-mRNA /GEN-NUCB2 / PROD-nucleobindin 2 / DB_XREF=g1:4826869 /
203675_at	UG-HS.3164 nucleobindin 2 /FL-gb:AF052542.1 gb:AF052643.1 gb:NA_065013.1 GEN=CPR8 /PROD=cell cycle
221156_x_at	gb:NM_004748.1 /DEF=Homo sapiens cell cycle progression 8 protein /FL=gb:AF011794.1 gb:NM_004748.1 progression 8 protein /DB_XREP=gi:4758047 /UG=Hs.283753 cell cycle progression 8 protein /DB_XREP=gi:4758047 /UG=Hs.283753 cell cycle progression 8 protein /PL=gb:AF011794.1 gb:NM_004748.1
207791 s at	gb:NM_004161.1 /DEF=Homo sapiens RAB1, member RAS oncogene family (RAB1), mRNA. /FEA=mRNA /GEN=RAB1 /PROD=RAB1,

	member RAS oncogene family /DB_XREF=gi:4758987 /UG-Hs.3642 RAB1, member RAS oncogene family /FL-gb:NM_004161.1 gb:MZ8209.4
	gb:NM. 004134.1 /DEF-Homo sapiens heat shock 70kD protein 9B (mortalln-2) (HSPA9B), mRNA. /FEM-mRNA /GEN-HSPA9B /PROD-heat shock 70kD protein 9B (mortalln-2) /DB_XREF-gi:4758569 /UG-Hs.3069 heat shock 70kD protein 9B (mortalln-2)
200692_s_at	/FL=gb:BC000478.1 gb:L15189.1 gb:NM_004134.1 gb:NM_014167.1 /DEF=Homo sapiens HSPC128 protein (HSPC128), mRNA. /FEA=mRNA /GEN=HSPC128 /PROD=HSPC128 protein
218936_s_at	/DB_XREF=gi:7661789 /UG=Hs.90527 HSPC128 protein /Firego:Arits17:1 95741517:1 95741517:1 95741518 /UB XREF=gi:7661789 /UB XREF=gi:76489138 /UB XREF=gi:76489
213931_at	/UG=HS.180919 inhibitor of DNA binding 2, dominant negative helix-loop-nellx protein. [JOS51320] /PEA-ERNA /GEN-LOC51320 /PROD-hypothetical protein [JOS51320] mRNN. /PEA-ERNA /GEN-LOC51320 /PROD-hypothetical protein // PEA-ERNA / PEA-ERNA / GEN-LOC51320 /PROD-hypothetical protein // PEA-ERNA / PEA-ERNA / GEN-LOC51320 /PROD-hypothetical protein // PEA-ERNA / PEA-ERNA / GEN-LOC51320 /PROD-hypothetical protein // PEA-ERNA / PEA
218247_s_at	DB_XREF=gi:7706165 /UG=HS.12830 hypothetical procesn /ringinal grandon friegosate for the feather feather for the feather feat
213935_at	/UG=Hs.184019 Homo sapiens clone 23551 mRNA sequence
208815_x_at	GD: ADUZ-920.1 / DER-Hamin Capitali 90093 heat shock 70kD protein 4 /FL=gb: AB023420.1 / PROD=apg-2 / DB XREF=g1:4579908 / UG=HS.90093 heat shock 70kD protein 4 /FL=gb: AB023420.1
218177_at	Consensus includes gb:AA293502 /FEA=EST / DB_AKGF=g1:1341030 / DB_AKGF=g1:1341030 / DB_AKGF=g1:1341030 / DB_AKGF=g1:1341030 / DBABARNA / GEN-GLUL / CHMP1.5 protein / FEA=mRNA / GEN-GLUL /
•	gb:NN 002065.1 /DEF-Homo sapiens glutamate-ammonia 1998se (glutamine synthase) /US-HS.170171 PROD-glutamate-ammonia ligase (glutamine synthase) /UB.XREF-gl:4504026 /UG=HS.170171
200648_s_at	glutamate-ammonia ingase (glutamine sylinase) /ri-glimate-ammonia ingase (glutamine sylinase) /ri-glimate-ammonia ingase (glutamine sylinase) /ri-glimate-ammonia ingase (glutamine sylinase) /ri-glimate (glutamine sylinase) /ri-glim
202843 at	FKUDEHLICVORGULAR GROCKLELT DIS AND STATE OF THE STATE OF
	Consensus includes gb:AV683882 /FEA=EST /DB XREF=gi:10285745 /DB XREF=est:AV683882 /CLUME=carcalcus /ug=ms:1/020
214578_sat	KNO-associated, College College Districtions appears natural resistance-associated macrophage protein 1 (SLC11A1) gene, Consensus includes gb: AF229163 /DEF=Homo sapiens natural resistance-associated macrophage protein 1 (SLC11A1) gene,
	complete cds, alternatively spliced; and nuclear LIM interactor-interacting ractor (NLT AT) year, Carrier and nuclear LIM interactor-interaction (NLT AT) year, 10257408 (UG=HS.182611 solute carrier family 11 (proton-coupled divalent metal ion transporters).
217473 x_at	member 1
	gb:NW_000328.1 /DEF=Homo sapiens retinitis pigmentosa GTPase regulator (KPGK), Makha: /fir-makha. pigmentosa GTPase regulator /DB_XREF=g1:4506580 /UG=Hs.153614 retinitis pigmentosa GTPase regulator /
207624_s.at	FL=gb:U5/629.1 gb:NM_000328.1 hb:NM_001453.1 /DEF=Homo sapiens zinc finger protein 198 (ZNF198), mRNA. /FEA=mRNA /GEN=ZNF198 /PROD=zinc finger protein
202778_s_at	198 /DB XREF=gi:4508010 /UG=Hs.109526 zinc finger protein 198 /FL=gb:AF035374.1 gb:AF060181.1 gb:NM_U03453.1
200653 s at	gb:MZ7319.1 /DEF-Human calmodulin mRNA, complete cds. /FEA-mKNA /FAOD-Calmodulin /DEF-Human calmodulin (phosphorylase kinase, delta) /FL=gb:MZ7319.1 gb:MX_006888.1
218627 at	gb:NM_018370.1 /DEF=Homo sapiens hypothetical protein FLJ11259 (FLJ11259), mkNA: /FEA=mNA: /GEN-ED1225, /LICOLDE PROTEIN PROTE
	mKNA. /red=mkNA /gen=arsva. in alpha 5) /FL=gb:BC002374
202059_s_at	gb: BCU13UUS.1 gb:NM U04204 gb:NM 024075.1 / DEF=Homo sapiens LENGS protein (LENGS), mRNR. / FEA=mRNR / GEN=LENGS / PROD=LENGS protein /
218132_s_at	od14f07.s1
203265_s_at	UG-Hs.75217 mitogen-activated protein kinase kinase 4 /FL=gb:NM_003010.1 gb:L36870.1 gb:U1//43.1
. 10000	gb:M97655.1 /DEE=Human 6-pyruvoy1tetranydropterin synthase (F13) maxii comprovitetranydropterin synthase (Pp. 1976) propeesin synthase (JDE_REFEG1:306438 /UG=HS.366 6-pyruvoy1tetranydropterin synthase (JDE_REFEG1:306438 /UG=HS.366 6-pyruvoy1tetranydropterin synthase (JDE_REFEG1:306438 /UG=HS.366 6-pyruvoy1tetranydropterin synthase (JDE_REFEG1:306438 /UG=HS.366 6-pyruvoy1tetranydropterin synthase
217591 at	:BF725121
222309_at	Consensus includes gb: AM972292 /FEA=EST /DB_XREF=gi:8162138 /DB_XREF=est:EST384381 /UG=Hs.292998 ESTs

	Hab:NW 005542.1 /DEF=Homo sapiens insulin induced gene 1 (INSIG1), mRNA. /FEA-mRNA (GEN=INSIG1 /	<u></u>
201627_s_at	1 / DB XREF=gi:5031800 /UG=Hs.56205 insulin induced gene 004544 /DEF=Homo sapiens BAC clone CTB-16284 from 4 /FEA=C	T
217249_x_at	UG=Hs.248095 Homo sapiens BAC clone CTB-10254 from 4 gb:NM_016509.1 /DEF=Homo sapiens C-type lectin-like receptor-2 (LOC51266), mRNA. /FEA-mRNA /GEN=LOC51266 / PROD=C-type lectin-like receptor-2 /DB_XREF=g1:7706060 /UG=Hs.114231 C-type lectin-like receptor-2 /FL=gb:AF124841.1	
220496_at	gb:NM_016509.1 Consensus includes gb:AL537042 /FEA=EST /DB_XREF=gi:12800535 /DB_XREF=est:AL537042 /CLONE=CSODF017XF17 (5 prime) /	
201096_s_at	UG=HS. 75290 ADP-ribosylation factor 4 (rh=gp:BcU03504.1 gb:mcorresponded ADP-ribosylation (3 prime) (Consensus includes gb:AL544094 (FER=EST) DB_XREF=gs:12876573 (DB_XREF=est:AL544094 (CLONE=CSODIO04YG20 (3 prime) (Consensus includes gb:AL544094 (FER=EST) DB_XREF=gs:12876573 (DB_XREF=est:AL544094 (CLONE=CSODIO04YG20 (3 prime) (Consensus includes gb:AL544094 (FER=EST) DB_XREF=gs:12876573 (DB_XREF=est:AL544094 (CLONE=CSODIO04YG20 (3 prime) (Consensus includes gb:AL544094 (FER=EST) (DB_XREF=gs:12876573 (DB_XREF=est:AL544094 (CLONE=CSODIO04YG20 (3 prime) (CONSENSUS INCLUDES gb:AL544094 (CLONE=CSODIO04YG20 (3 prime) (CLO	· -
201580_s_at	FEST /DB_XREF=91:10836094 /DB_XREF=est:AV726673 /CLONE=HTCBGC	
201109_s_at	a	
203080_s_at	domain, 2B /FL-gb:ABU34255.1 gb:Nm_1534001 gb:Nm_0069681.1 /DFF=Homo sapiens stress 70 protein chaperone, microscome-associated, 60kD (STCH), mRNA. /FEA=mRNA / GRN=STCH /PROD=stress 70 protein chaperone, microscome-associated, 60kD /DB_XREF=gi:5902125 /UG=Hs.288799 stress 70	
202558_s_at	some-associated, 60 sapiens solute camenna /GEN=SLC9A3R1	Ī
201349_at	<pre>factor 1 /DB_XREF=g1:4759139 /UG=Hs.184776 solute carrier iamily 9 (solutumyAragan exchaller);</pre>	· T
220603_s_at	gn:www_loss:1 / Dzf.nown orgin: 32 NREF=gi:8922916 /UG-HS.33368 hypothetical protein FLJ1175 /FL=gb:NW_018349.1 PROD=hypothetical protein FLJ1175 /DB_XREF=gi:8922916 /UG-HS.33368 hypothetical protein FLJ1175 /FL=gb:NW_018349.1	<u> </u>
35820_at	Cluster Incl. X62078:H.sapiens mRNA for GM2 activator protein /cds=UNKNONN /gb=X62078 /gi=313158 /ug=Hs.69743 /len=2436	Τ
	gb:NN_003292.1 /DEF=Homo sapiens translocated promoter region (to activated MrP. oncogene) (1FF);	1
201731_s_at	promoter region (to activated has ancogene, free some complementing rodent repair deficiency, complementation group 5 gb:NN_000123.1 /DEF=Homo sapiens excision repair cross-complementing rodent repair /FEA=mRNA /GEN=ERCC5 /	
	complementation group 6 (Lockayne Symbolicy) (Lockayne Symbolicy) (Lockayne Symbolicy) (Lockayne Symbolicy) (Lockayne Symbolicy) (Keroderma pigmentosum, complementation group 6	
202414_at	FL=gb:D16305.1 gb:L20046.1 gb:NM_U00123.1 gb:D50402.1 /DEF=Human mRNA for NRAWF1, complete cds. /FEA=mRNA / PROD=Nramp / DB_XREF=gi:1000996 /UG=Hs.182611 solute	
210422_x_at	carrier family 11 (proton-coupled divalent metal ion transporters), memor since since and since since since since and since si	· T
58900 at		i
	arbonydrate-recognition dumain; rectin, diperaterical dependent, carbohydrate-recognition domain) lectin, (calcium dependent, carbohydrate-recognition domain)	- .
219890_at	Tectin, superjeanily memoer 3 (ringuismicrozoni garning) (REDI), mRNA. (FERA-mRNA (GENEEND) /FROD=EH domain containing 1 / gb:NN 006795.1 (DEPENEEND) (PROD=EH domain containing 1 / Flach: RP0999011.1 db:NN 006795.1	
208112_x_at	DB XXEF=91.3003000 /OG-BS.13317 20 CONTROLL DESCRIPTION OF THE STATE O	-1
218655_S_at	robengybourertest process. (FER-EST /DB_XREF=g1:795120 /DB_XREF=est:yf61e05.s1 /CLONE=IMAGE:26577 / Conseasus includes gb:R7512064 (FER-EST /DB_XREF=g1:795120 /DB_XREF=est:yf61e05.s1 /CLONE=IMAGE:26577 / Inc.Hs. 16449 cell division cycle 42 (GTP-binding protein, 25kD)	$\overline{}$
201898_s_at	Consensus includes gb:AI126625 /FEA=EST /DB_XREF=gi:3595139 /DB_XREF=st:qd83a12.x1 /CLONE=IMAGE:1/30002 / UG=Hs.80612 ubiquitin-conjugating enzyme EZA (RAD6 homolog) /FL=gb:M74524.1 gb:NM_003336.1	7
		•

	gb:U81501.1 /DER-Buman lipoxin A4 receptor mRNA, complete cds. /FEA=mRNA /PROD=lipoxin A4 receptor /DB_XREF-gi:1916075 / mc=Hs. 99855 formyl peptide receptor-like 1 /FL=gb:M76672.1 gb:M88107.1 gb:U81501.1 gb:AF054013.1
210773_s_at	gb:NM_001462.1
32069 <u>a</u> t	Cluster Incl. AB01415:Homo sapiens mayn lor Arrocis grows, Ossay. ug=Hs.155972 /len=3319
121_at	X69699 /FEATURE / DEFINITION-HSPAX8A H. Sapiens Pax8 mRNA commlete cds. /FEA-mRNA /GEN-FALDH /PROD-fatty
	gb:L47162.1 /DEF=Human ratty aldenyde denydrogenase (ranka) maken a family, member A2 /Fi-gb:L47162.1 aldehyde denydrogenase /DE,XREF=gi:1082035 /UG=Hs.159608 aldehyde dehydrogenase 3 family, member A2 /Fi-gb:L47162.1
202053_s_at	gb:U4bb89.1 gb:NM_U0U364.1 Consensus includes gb:AM167793 /FEA=EST /DB_XREF=gi:6399401 /DB_XREF=est:xg56d07.xl /CLONE=IMAGE:2632333 /UG=Hs.164036
212335_at	Homo sapiens AKAP350C mRNA sequence, alrernatively spliced Ch. Ni 678 1 / NPR-Human T-cell receptor gamma chain VJCI-CIII region mRNA, complete cds. /FEA-mRNA /GEN-TCRG /
209813_x_at	DE XREF=gi:339399 /UG=Hs.112259 T cell receptor gamma locus /PL=gb:M16768.1 gb:AF151103.1
	sapiens interferon-induced protein With retraticopeptide repeats 4 / DB XRE-gi:450456 / UG=Hs.181874 on-induced protein withtetratricopeptide repeats 4 / DB XRE-gi:450456 / UG=Hs.181874
	interferon-induced protein with tetratricopeptide repeats 4 /FL=gb.BC001383.1 gb:BC0049//.1 gb:D2513.1 gb:Account.
204747_at	EBIARUSES N.1 1987 N. 19126544 (FER-EST /DB XREF=g1:5662508 /DB XREF=est:wo46c12.x1 /CLONE-IMAGE:2458390 /UG=HE.303154
212223 at	DODGEVE DYCKETH 3
	Consensus.includes gb:N35408 /FEA-EST /DB_XREF=gi:1157550 /DB_XREF=est:yy33103.sl /LLUNE=LEAGE:2/3033 /OG=LS:333350 / OG=LS:333350 / OG=LS:33350 / OG=LS:3350 /
218880_at	hypothetical protein Fileson (Firesonal Proteins), mana, /FEA-mana /GEN-LPINZ /PROD-lipin 2 /DB_XREF=gi:7662021./
202460_s_at	UG=Hs.166318 lipin 2 /FL=gb:D87436.1 gb:NM_014646.1
20790 0 00) end /crone-mass-joss / crone-
30100_8_84	sapiens phosph
205281 s at	transcript Variant 1, mkWa. / Linguistry (Single of Control of Con
	gb:NM_021242.1 /DEF=Homo sapiens hypothetical protein STRAIT11499 (STRAIT11499), mRNA. /FEA=mRNA./GEN=3.FALIII.
218251_at	PROD=mypochetical protein Signification / DE_Man Series / DE_XREF=est.ap29d10.x1 /CLONE=IMAGE:1956787 /
212481_s_at	UG=Hs.250641 tropomyosin 4
213292 s_at	Consensus includes girlandoring the consensus from the consensus formulate cds. /
٠	gi:13529208 /UG=Hs.6523 chromosome 1 open
221497_x_at	FL=gb:AF229245.1 gb:AF277176.1 gb:NM 022051.1 gb:BC005369.1
212185 x at	Consensus includes gb:NM_U05953.1 / DBF-namb Saprems includes 28 /FL-gb:NM_U05953.1 PROD-metallothionein 2A /DB_XRBF=g1:5174763 /UG-H3.118786 metallothionein 2A /FL-gb:NM_U05953.1
	L157398 /DEF=Human DNA sequence from clone RP11-5bH/ on chromosome 10. Concarno 1207 100 100 ne nebulette protéin (NEBL, actin-binding Z-disc protéin) /FEA=mRNA_1 /DB_XREF=gi:10045326 /
203961_at	UG=Hs.5025.nebulette /FL=gb:NM_006393.1
1	gb:Z24/27.1 / DER=H.sapiens tropomyosin isotorm mada, complete CDS. /FEA-mada / Fine-Logonyosin isotorm mada /
210986_S_ac	Human mRNA
41386_1_at	len=6121 Concensis includes db: BP978611 /FEA=EST /DB_XREF=gi:12345826 /DB_XREF=est:602149061F1 /CLONE=IMAGE:4307822 /
201874_at	UG=HS.14891 hypothetical protein FLJ21047 /FL=gb:NM 024569.1 UG=HS.14891 hypothetical protein FJZ1047 /FL=gb:NM 024569.1
211781_x_at	gb:BC000164.1 / DBF-BAMBO Sapitus, Civil 2000 10 10 10 10 10 10 10 10 10 10 10 10
14 p 074000	
20-8-12-0-10	

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	sapiens PC326 protein (PC326), mRNA. /FEA-mRNA /GEN
217908_s_at	UG-HS.279882 PC32b procein /ri-gd:Allicology. gurricology. School Control Cont
202427_s_at	protein /DB XREF=g1:7661601 /UG=Hs.76285 DKFZF5564B167 protein /FL=gD:ALLIUZ9:.1 gD:NR_UL19415.1.
213716_s_at	Home sapiens secreted and transmembrane 1 (SECTM1), mRNA Home sapiens secreted and transmembrane 1, 24064 Home
212550 at	Consensus includes gb:AI149535 /FEA=EST / DB_XKE=g1:0:0004 / DS_XKET=G1:0:0004 / DS_XK
11101 r at	Cluster Incl. X57348:H.sapiens mRNA (clone 9112) /cds=(165,911) /gb=X57348 /gi=23939 /ug=Hs.184510 /len=1407
75-7-09000	protein (DKFZP434P1750), mRNA. /FEA=mRNA /GEN=DKFZF434F1 /HG=Hs. 7274 DKFZP434P1750 protein /FL=gb:NM_015527.1
220947_s_ar	PROD=DAK244441700 process / PER=mRNR / Pp:NNL-002387.2 / DEF=Homo sapiens Wiskott-Aldrich syndrome protein interacting protein (WASPIP), mRNA. / FER=mRNR / Gran-wacpip / PROD=WASP-interacting protein /DB_XREF=gi:8400739 /UG=Hs.24143 Wiskott-Aldrich syndrome protein
202665_s_at	interacting protein /FL=gb:NM 003387.2
	sapiens retinoid x receptor interacting protein /DE_XREF=gi:7706482 /UG=Hs.7889 retinoid x rece
220746_s_at	FL=gb:AF113538.1 gb:NM_016290.1
218078_s_at	gbink_utop96.1 / Ubr=ncum Saptum Saptum /FL=gb:Ar247703.1 gb:NM_016598.1 DB_XREF=gi:7706132 /UG=Hs.14896 DHHC1 protein /FL=gb:Ar247703.1 gb:NM_016598.1
76 2 PC/200	gb:NM_016227.1 /DEF=Homo sapiens membrane protein CH1 /FL=gb:NF097535.1 gb:NM_016227.1
	ns tumor protein p53-binding protein, 2 (TP53BP2), mWWA. protein, 2 /DB_XREF=g1:4885642 /UG=HS:44585 tumor protei
203120_at	/FEL=gb:U58334.1 gb:NM_005426.1
1000	Consensus includes gb:T79953 /PEA=EST /DB_XREF=G1:b98402 /DB_AKEF=ESL:ydd0J11:31 /CDGAL FT180040 gene product /PF=ch:D2539.1 db:NM 014656.1
203143_S_ac	
41387_r_at	len=6121
202286_s_at	Conseisus includes 92.00.22. / UG-HS.13582 tumor-associated calcium signal transducer 2 / FL-go:NM U0233.1 / Clone 05516 / FRA-mRNA / DB_XREF-gi:182893 / UG-HS.23582 tumor-associated calcium signal transducer 2 / FL-go:NM U0233.1
4. 053000	gb:NM_005080.1 /DEF=Homo sapiens X-box binding protein 1 (XBF1), makk: /FEA-makk / Pan-A21.1 (1905080.1 hn YRFP-c: 4827057 /UG-HS.149923 X-box binding protein 1 /FL=gb:BC000938.1 gb:N31627.1 gb:NM_005080.1
2000/0-01	gb:062858.1 /DEF=Human interleukin-13 receptor mRNA, complete cds. /FEA=mRNA /PROD=interleukin-13 receptor /
211612_s_at	DB_XRRF=01:1658578 /FL=gb:1628283
203233 at	gb:NM_000418.1 / DEF=Homo Sapiens incerteur. 1000418.1 precursor /DB_XREF=g1:4557668 /UG=Hs.75545 interleukin 4 receptor /FL=gp:NM_000418.1
	gb:NM_000945.1 /DEP=Homo sapiens protein phosphatase 3 (formerly AB), regulatory submitt 2 (2007) - 2 (2007) (1007
	regulatorysubunit B (19kD), alpha isoform (calcineurin B, type I) /DB_XREF=gi:4506024 /UG=HB.//8340 process process process regulatorysubunit B (19kD), alpha isoform (calcineurin B, type I) /FL=gb:M30773.1.gb:NM_000945.1
204507_s_at	Hotmerly 28, Legulatory success interhondrial solute carrier (LOC51312), mRNA. /FEA-mRNA /GEN-LOC51312 /PROD-hypothetical gb:NM_018579.1 /DEF-Homo sapiens mitochondrial solute carrier (LOC51312), mRNA. /FEA-mRNA /GEN-LOC51312 /PROD-hypothetical gb:NM_018579.1 /DEF-Homo sapiens mitochondrial solute carrier /FIE-dp:NPE155660.1 gb:NF116630.1
)18136 c at	protein PRO1278 /DB XXEE=g1:83/40// /UG=ns.Judgy militaring for the last exm of the last exm of
	Consensus includes gb:AL121900 /DEF=Human DNA sequence from clone RP11-3/905 on cnromosome for a novel protein
201582_at	B / KL=gD: ECUDATUR. 1 go. AL. 2013. Complete cds. / FEA-mRNA / PROD=hSIAH2 / DB_XREF=g1:2673967 / UG=Hs.20191 seven in absential db:U76248.1 / DEF=Human hSIAH2 mRNA, complete cds. / FEA-mRNA / PROD=hSIAH2 / DB_XREF=g1:2673967 / UG=Hs.20191 seven in absential
209339_at	[Drosophila] homolog 2 /FL=gb:U76248.1 gb:NM_005067.1 [Drosophila] nRNA. /FEA=mRNA /GEN=ADD1 /FROD=adducin 1 https://doi.org/10.1007/11.0 1001119 2 /DPPEHomo sapiens adducin 1 (alpha) (ADD1), transcript variant 1, mRNA. /FEA=mRNA /GEN=ADD1 /FROD=adducin 1
208030_s_at	gbing_orly.

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Tabelle 4: Gene aus Clusteranalyse 4

Affymetrix interne	Beschreibung der Sequenz in der Genebank Datenbank
Bezeichnung	
	includes gb:AI
221798_x_at	profess S. A. Arrenders National National National Property of Professional Property of Professional Professi
	gb:NM_01/20.1 / Lbr -no.nc suprems in no.1 / 10.000 / 10.
218231_at	kinase /DB_XREF=g1:8923736 /UG=HS./UJS N-ACELY1914COSAMALE ALTOSOFT / CLONE=INAGE:2073430 /UG=HS.179526
	Consensus incrudes garaticode and a consensus incrudes garaticode and a consensus incrudes a consensus incredible and a consensus
201009_s_at	by 1,25-dihydroxyvitamin D-3 /FL=gb:NM 006472.1 gb:S73591.1 by 1,25-dihydroxyvitamin D-3 /FL=gb:NM 006472.1 gb:S73591.1
	Consensus includes gb:AI583173 /FRA=EST / DE_AAKFFFgl:4303070 / DE_AAKFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF
212998 x at:	histocompatibility complex, class II, DQ beta 1
	Consensus includes gb:BG491844 /FEA=EST /DB_XREF=g1:13455550 /Jb_ArreF=esc:00255555 /
1	
201464_X_ac	Consensus includes gb:AK025577.1 /DEF=Homo sapiens cDNA: FLJ22924 fis, clone KAT06977, highly similar to include
	fibroblast
214687_x_at	A=mkNA / DB_Arct -gittotofa
,	gb:NM_014020.1 / DEF=Homo sapiens into process (2007) and (2007) a
220532_s_at	LR8 protein /FL=gb:AF115384.1 gb:NM_014020.1
	'n,
205987 at	lypeptide / DB XREF=g1:4502646 /UG=Hs.1311 CD1C antigen, c polypeptide /FL=gb:M28827.1 gb:NM_001765.1 DB XREF=g1:4502646 /UG=Hs.1311 CD1C antigen, c polypeptide /FL=gb:M28827.1 gb:NM_001765.1
	factor 11B (FEA11B), unders. / FEA-11B /FL=gb: AF093670.
202658 at	gb:NM_003846.1
	gb:NM_014748.1 /DEF=Homo sapiens KIAA0064 gene product (KIAA0064), haden (Kianoca)
200991 c at	- 33
11	gb:AF267856.1 /DEF=Homo sapiens HT033 mRNA, complete cds. /FEA=mkNA./FKOD=n1033 /DD_1200
	thetical protein
209007_s_at	dd465NZ4.2.1
213969_x_at	ribosomal protein L29
1	Consensus includes gb:AI004246 /FEA=ESI /DB_AREF-91:3213/30 /22315
200094_s_at	gb:BC004954.1 / DEF=Homo sapiens, clone MGC:10897, mRNA, complete cds. /FEA=mRNA / PROD=Unknown (protein 104 mark) gb:BC004954.1 jb:MM_000977.1
208929_x_at	'UG=Hs.180842 ribosomal protein uto

	sapiens adrenergic, beta-2-, receptor, surface (ADRB2), m
206170 at	/PROD=adrenergic, beta-2-, heceptor, surface /DB XREF=qi:13162366 /UG=Hs.2551 adrenergic, beta-2-, receptor, surface /FL=gb:NM_000024.2 gb:M15169.1
:	mRNA; cDNA DKFZp564E2282 (from cl
	/UDB_XKKEF=g1:4084109 /OG-AB-109093 ALTALISTS FINES FINES FOR A THIOLOGY INTERPORT OF THE TRANSF:2492333 CONSENSUS INCludes gb:AI972144 /FEA=EST /DB_XREF=g1:5768970 /DB_XREF=est:wr63b03.x1 /CLONE=IMAGE:2492333 /UG-Hs.75860 hydroxyacyl-Coenzyme A chydrogenase3-ketcacyl-Coenzyme A thiolaseenoyl-Coenzyme A hydratase /UG-Hs.75860 hydroxyacyl-Coenzyme A chydrogenase3-ketcacyl-Coenzyme A thiolaseenoyl-Coenzyme A hydratase
208630_at	712
214058_ac	/OG=HS.9213/ V-myc avian myelocycomecosis vira: possible aviant part / October 17030g05.x1 /CLONE=IMAGE:3700017 / Consens includes gb:RF2224 /FEA=EST /DB_XREF=gi:1130401 /DB_XREF=est:7q30g05.x1 /CLONE=IMAGE:3700017 / CONSENS includes gb:RF2324 /FEA=BST /DB_XREF=gi:1130401 /DB_XREF=est:7q30g05.x1 /CLONE=IMAGE:3700017 / CONSENS includes gb:RF2-gb:RF201:1130401 /DB_XREF=est:7q30g05.x1 /CLONE=IMAGE:3700017 / CONSENS includes gb:RF2-gb:RF201:1130401 /DB_XREF=est:7q30g05.x1 /CLONE=IMAGE:3700017 / CONSENS includes gb:RF201:1130401 /DB_XREF=est:7q30g05.x1 /CLONE=IMAGE:3700017 / CONSENS includes gb:RF201:1130401 /DB_XREF=est:7q30g05.x1 /CLONE=IMAGE:3700017 / CONSENS includes gb:RF201:1130401 / DB_XREF=est:7q30g05.x1 /CLONE=IMAGE:3700017 / CONSENS includes gb:RF201:1130401 / DB_XREF=est:7q30g05.x1 / CLONE=IMAGE:3700017 / CONSENS includes gb:RF201:1130401 / DB_XREF=est:7q30g05.x1 / CLONE=IMAGE:3700017 / CONSENS includes gb:RF201:1130401 / DB_XREF=est:7q30g05.x1 / CLONE=IMAGE:3700017 / C
212179 at	02 /DB_XREF=est:au83a02.x1 /CLONE=IMAGE:2782826
	gb:BC000533.1 /DEF=Homo sapiens, Similar to eukaryotic translation initiation factor 3, subunit 8 (110KD), clone MGC:8693,
210949_s_at	mRNA, complete cds. /FEA-mRNA, /PROD=Similar to eukaryotic translation initiationfactor 3, subunit 8 (110kD) /FI-29b=BC006533.1 // NB_XREF=gi:12653522 /UG=Hs.4835 eukaryotic translation initiation factor 3, subunit 8 (110kD) /FI-29b=BC006533.1
	Consensus includes gb:AV700415 /FEA=EST /DB_XREF=gi:10302386 /DB_XREF=est:AV700415 /CLONE=GKCDGAU4 /UG=ns.µz303 Lyz0- Therical protein
217817 at	Consensus includes gb:BE891920 /FEA=EST /DB_XREF=gi:10351728 /DB_XREF=est:601435490F1 /CLONE=IMAGE:3920590 /UG=Hs.323342 actin related protein 23 complex, subunit 4 (20 kD) /FL=gb:AF006087.1 gb:NM_005718.1
216041 × at	Consensus includes gb:AKO23348.1 /DEF=Homo sapiens cDNA FLJ13286 fis, clone OVARC1001154, highly similar to Homo sapiens clone 24720 epithelin 1 and 2 mRNA. /FEA=mRNA /DB_XREF=gi:10435243 /UG=Hs.180577 granulin
202687 s at	gb:U57059.1 /DEF-Homo sapiens Apo-2 ligand mRNA, complete cds. /FEA-mRNA /PROD-Apo-2 ligand /DB_XREF=gi:1346207 /UG=Hs.83429 tumor necrosis factor (ligand) superfamily, member 10 /FL-gb:U37518.1 gb:U57059.1 gb:NM_003810.1
215230 x at	/CLONE=IMAGE:1156734
	gb:NM_003752.2 /DEF=Homo sapiens eukaryotic translation initiation factor 3, subunit 8 (110kD) (EIF3S8), mkWA. /Fera=mrNA /GEN=EIF3S8 /PROD=eukaryotic translation initiation factor 3, subunit 8 (110kD) /DB_XREF=g1:5579457 ////////////////////////////////////
2000# 1_X_aL	gb:BC005876.1 /DEF=Homo sapiens, ATPase, H+ transporting, lysosomal (vacuolar proton pump) 21kD, clone MGG:4498, mRNA,
200078_s_at	~
	gb:AF302110.1 /DEF=Homo sapiens alpha-aminoadipic semialdehyde dehydrogenase-phosphopanternelnyl transferase complete cds. /FEA=mRNA /PROD=alpha-aminoadipic semialdehydedehydrogenase-phosphopantetheinyl transferase //DB_XREF=gi:11120434 /UG=HS.64595 aminoadipate-semialdehyde dehydrogenase-phosphopantetheinyl transferase
202169_s_at	/FL-gb:AF302110.1 gb:AF1369/8.1 gb:AF151838.1 gb:AF15103/.1 gb:NN_012423.1 gb:NN_072801.1 /DEF=Homo sapiens proteasome (prosome, macropain) subunit, beta type, 10 (PSMB10), mRNA. /FEA=mRNA
202659 at	יייייייייייייייייייייייייייייייייייייי
	/DEF=Homo sapien
221488_s_at	protein CUTA /FL=gb:AF230924.1
221622_s_at	erized hypothalamus protein HT007 /FL-gb:AF24

	gb:NM_002939.1 / DEF=Homo sapiens ribonucleaseangiogenin inhibitor (RNH), mRNA. / FEA=mKNA / GEN=KNH
206050_s_at	39
12100 at	Consensus includes gb:AL566962 /FEA=EST /DB_XREF=gi:12919867 /DB_XREF=est:AL566962 /CLONE=CSUDFU26YHU8 (3 prime) ///TG=Hs 284281 Human outative ribosomal protein Sl mRNA
	Consensus includes gb:BF965566 /FEA=EST /DB_XREF=gi:12332781 /DB_XREF=est:602277032F1 /CLONE=IMAGE:4364790
201861_S_at	Oug=Rs.320139 leucine fich repeat (in Fill) interacting process. Consensus includes gb:AL137753.1 / DEF=Homo sapiens mRNA; cDNA DKFZp434K1412 (from clone DKFZp434K1412).
212795_at	/FEA=mRNA /DB_XREF=g1:6808455 /UG=HS.1Z144 KIAALU33 protein Consensus includes qb:AA861608 /FEA=EST /DB_XREF=g1:2953748 /DB_XREF=est:aK34e01.s1 /CLONE=IMAGE:1407864
213574_s_at	/UG=Hs.180446 karyopherin (importin) beta 1
	Consensus includes gb:AK023063.1 /DEF=Homo sapiens cDNA FLJ13001 fis, clone NIZKP3000341, nignry similar of nomb Sapiens mitochondrial inner membrane preprotein translocase Tim17a mRNA, nuclear gene encoding mitochondrial protein.
215171_s_at	/FEA=mRNA /DB_XREF=gi:10434808 /UG=Hs.20716 translocase of inner mitochondrial membrane 17 (yeast) homolog(A
210137's at	gb:BC001286.1 /DEF=Homo sapiens, Similar to dCMP deaminase, clone MGC:5160, mRNA, complete Cds. /FEA=mkNA /PROD=Similar to dCMP deaminase /DB_XREF=gi:12654884 /UG=Hs.76894 dCMP deaminase /FL=gb:BC001286.1
212943 at	Consensus includes gb:AB011100.2 /DEF=Homo sapiens mRNA for KIAA0528 protein; partial cds. /FEA=mRNA /GEN=KIAA0528 //PRON=KTAA0658 protein /DB XREF=gi:6683714 /UG=Hs.30656 KIAA0528 gene product
	gb:AF180476.1 /DFF=Homo sapiens CALIFP (CALIF) mRNA, complete cds. /FEA=mRNA /GEN=CALIF /PROD=CALIFP /DB_XREF=gi:6856208
202164_s_at	/UG=HS.Zb/U3 CCR4-NOT Cranscription complex, smounted by Fig. and System Systems of PROD=D45SKP2-like protein mRNA. complete cds. /FEA=mRNA /PROD=D45SKP2-like protein
209005_at	gb:AFIJ/323.1 / DEF=num Sapieus P-300.2 12.0 process from 5 /FL-gb:AF199420.1 gb:AF142481.1 gb:AF157323.1 // DB XREF=gi:7688696 /UG=HS.5548 f-box and leucine-rich repeat protein 5 /FL-gb:AF199420.1 gb:AF142481.1 gb:AF157323.1
	no sapiens
202899 s at	INGINITIESELIMETICA) / DD_AKEF-91:4300000 / OG-MS:401300 BF-10130
ή .	me subuni
208777 g at	
ш	gb:NM_005479.1 /DEF=Homo sapiens frequently rearranged in advanced T-cell lymphomas (FRAT1), mRNA. /FEA=mRNA
	_
219889_at	
	gd:NM_U0U0253.1 / LEF=Homo Sagrems mentioning Cycosolic factor I (4//L) discuss section 1 (4//L) forther menny
100000	FERA-MRNA /GEN-NCF1 /PROD-meutrophil cytosolic factor 1 /DB_XREF-gi:4557784 /UG=HS.1583 neutrophil cytosolic factor 1 / DB_XREF-gi:4557784 /UG=HS.1583 neutrophil cytosolic factor 1 / Arterior cytosolic factor 1 / FIL-ab:BC002816.1 db:AF330627.1 db:M55067.1 db:M25665.1 db:NM_000265.1
23	Consensus includes gb:AF091078.1 /DEF=Homo sapiens clone 559 unknown mRNA, complete sequence. /FEA=mRNA
217106_x_at	/PROD=unknown /DB XREF=g1:3859993 /UG=HS.145519 purative dimetily indemostraction from /PRA=CDS /GEN=ALDH1
	1, soluble /DB_XREF=gi:4502030 /UG=Hs.76392 aldehyde dehydrogenase
212224_at	
204125 at	gb:NM_016013.1 /DEF=Homo sapiens CGI-65 protein (LOC51103), mkNa. /FEA=mkNa /GEN=LOC51103 /FNC5-CGI 52 Protein /FL=gb:BC000780.1 gb:AF151823.1 gb:NM_016013.1
	gb:NM_018235.1 /DEF=Homo sapiens hypothetical protein FLJ10830 (FLJ10830), mRNA. /FEA=mRNA /GEN=FLJ10830
217752_s_at	/PROD=hypothetical protein FLJ10830 /DB_XREF=gi:8922698 /UG=Hs.273230 hypothetical protein FLJ10830 /FL=gb:BC001375.1 gb:BC003176.1 gb:NM_018235.1 ·
.221087_s_at	gb:NM_014349.1 /DEF=Homo sapiens apolipoprotein L, 3 (APOL3), mRNA. /FEA=mRNA /GEN=APOL3 /PROD=apolipoprotein L, 3

5 :

	/DB_XREF=g1:7656972 /UG=Hs.241535 apolipoprotein L, 3 /FL=gb:AY014905.1 gb:AF070675.1 gb:NM_014349.1
	gb:NN_018229.1 /DEF=Homo sapiens hypothetical protein FLJ10813 (FLJ10813), mRNA. /FEA=mRNA /GEN=FLJ10813 /PED=PLJ10813 /PED=PLJ10813 /DB_XREF=gi:8922687 /UG=Hs.106210 hypothetical protein FLJ10813 /PED=PLJ10813 /
218139_s_ac	/filego:Anisocos.r go:nw_vices.r. foresmisus includes gb:AIG6544 (FRA-EST /DB_XREF=est:tu38g02.x1 /CLONE=IMAGE:2253362
7138/8 <u>ac</u>	JOS-DB. 233009 Nerg process The last the last protein FLJ21432 (FLJ21432), mRNA. /FEA=mRNA /GEN=FLJ21432 (FLJ21432), mRNA. /FEA=mRNA /GEN=FLJ21432 (PRJ21432), mRNA. /FEA=mRNA /GEN=FLJ21432 /DB_XREF=gi:13375714 /UG=HS.11641 hypothetical protein FLJ21432
201340_ac	results includes gb:ALTSTANCE for a protein kinase, the COLITAL gene for collagen type XVII alpha 1 (BP180), contains the gene KIAA0204 (HSLK) for a protein kinase, the COLITAL gene for collagen type XVII alpha 1 (BP180), ESTs and GSSS /FEA-mRNA_2 /DB_XREF=gi:8573811 /UG=Hs.105751 Ste20-related serinethreonine kinase
206874_s_at	/FL-gb:D86959.1 gb:NM_014720.1 Consensus includes gb:NM_004251.1 /DEF=Homo sapiens RAB9, member RAS oncogene family (RAB9), mRNA. /FEA=CDS /GEN=RAB9
221808 at	member RAS oncogene tam.ly
	gb:BC005297.1 /DEF=Homo sapiens, Similar to kynurenine 3-monooxygenase (kynurenine 3-hydroxylase), clone MGC:12362, mkNA, complete cds. /FEA-mkNA /PROD=Similar to kynurenine 3-monooxygenase(kynurenine 3-hydroxylase) /DE_XREF=gi:13529016
777738"S"GC	
201386_s_at	פראטייי מחלטינייי יוויים ביים ביים ביים ביים ביים ביים
•	gb:NM_014751.1 /DEF=Homo sapiens KIAA0429 gene product (KIAA0429), mRNA. /FEA=mRNA /GEN=AIAA0429 /ENCD=AIAA0422 gene product
203037_s_at'	Is.77694
218356_at	sapiens cell division protein FtsJ (FUHL), mkNA. /FEA=mkNA /GENFOUL /FNO-Cell division protein FtsJ /FL=gb:AF093415.1 gb:NM_013393.1
	gb:NM_014473.1 /DEF=Homo sapiens putative dimethyladenosine transferase (HSA9761), mkNA /reA=mkNA /GEN3701 / /PROD=putative dimethyladenosine transferase /DB_XREF=gi:7657197 /UG=Hs.125819 putative dimethyladenosine transferase
204405_x_ar	/rL=gp:Ario1214/.1 gp:um_0144/0.1 qb:NM_007062.1 /DEF=Homo sapiens nuclear phosphoprotein similar to S. cerevisiae PWP1 (PWP1), mRNA. /FEA=mRNA /GEN=PWP1
201608 s at	
14	gb:NM_015380.1 /DEF=Homo sapiens CGI-51 protein (CGI-51), mRNA. /FEA=mRNA /GEN=CGI-51 /PROD=CGI-51 protein
201569_s_at	.
222244 s at	00749.1 /DEF=Homo sapiens cDNA FLJ20742 fis, clone HEP06891. /FEA=m. protein FLJ20618
	39.1 /DEF=Homo H (Asp-Glu-Ala-
201241 at	4939.1
220731_s_at	sapiens hypothetical protein FLJ10420 (FLJ10420), mRNA./FEA in FLJ10420 /DB_XREF=gi:8922415 /UG=Hs.289087 hypothetical pr
208799 at	sapiens, proteasome (prosome, macropain) subunit, beta type, 5, cione Mac:2115, mana,
22-22-02	

	/FEA=mRNA /PROD=proteasome (prosome, macropain) subunit, betatype, 5 /DB_XREF=gi:13278740 /UG=Hs.78596 proteasome
	(prosome, macropain) subunit, beta type, 5 /FL=gb:BC004146.1 gb:NM_002797.1 gb:D29011.1
221718_s_at	gb:M90360.1 /DEF=Human type II cAMP-dependent protein kinase (Ht31) mRNA, complete cds. /FEA=CDS /GEN=Ht31 /PROD=protein kinase /DB_XREF=gi:184434 /FL=gb:M90360.1
208662_s_at	Consensus includes gb:AI885338 /FEA=EST /DB_XREF=g1:5590502 /DB_XREF=est:w192e09.x1 /CLONE=IMAGE:2432392 /UG=Hs.118174 tetratricopeptide repeat domain 3 /FL=gb:D84294.1
202118 s at	Consensus includes gb:AA541758 /FEA=EST /DB_XREF=g1:2288192 /DB_XREF=est:ni87d05.s1 /CLONE=IMAGE:983817 /UG=Hs.14158 copine III /FL=gb:AB014536.1 gb:NM_003909.1 gb:AF077226.2
	gb:NN_004551.1 /DEF=Homo sapiens NADH dehydrogenase (ubiquinone) Fe-S protein 3 (30kD) (NADH-coenzyme Q reductase) (NDUFS3), mRNA. /FEA=mRNA /GEN=NDUFS3 /PROD=NADH dehydrogenase (ubiquinone) Fe-S protein 3(30kD) (NADH-coenzyme Q reductase)
201740 at	Lase, /DB_XEF=g1:4758787 /UG=Hs.5273 NADH dehydrogenase (ubiquinone) Fe-S protein 3 (30kD) (NADH-coenzyme Q reductase) /FE=qb:BC000617.1 qb:AF067139.1 qb:NM_004551.1 gb:AF100743.1
	Consensus includes gb:AF090094.1 /DEF=Homo sapiens clone IMAGE 172979. /FEA=mRNA /DB_XREF=gi:4063629 /UG=Hs.125078 ornithine decarboxylase antizyme 1
212796 s at	35
210561 s at	5564B0482); complete cds. /FEA=me 187991 DKFZP564A122 protein
	sapiens =Hs.2465
. 44 05200	gb:NM_006526.1 /DEF=Homo sapiens zinc finger protein 217 (ZNF217), mRNA. /FEA=mRNA /GEN=ZNF217 /PROD=zinc finger protein 217 /PRA=mRNA /GEN=ZNF217 /PROD=zinc finger protein 217 /PR=ah:AF041259 1 ah:NM 006526.1
211036 x at	יות חזוי
TE 2 03040C	sapiens protein k
64064_at	Cluster Incl. A1435089:th95b11.x1 Homo sapiens CDNA, 3 end /clone=IMAGE-2126397 /clone_end=3 /gb=A1435089 /gi=4300737 /ug=Hs.26194 /len=891
201515_s_at	
219093_at	gb:NN_017933.1 /DEF=Homo sapiens hypothetical protein FLJ20701 (FLJ20701), mRNA. /FEA=mRNA /GEN=FLJ20701 /PROD=hypothetical protein FLJ20701 /FL=gb;NM_017933.1
221739_at ·	Consensus includes gb:AL524093 /FEA=EST /DB_XREF=gi:12787586 /DB_XREF=est:AL524093 /CLONE=CSODC002XI04 (5 prime) /VG=Hs.10927 hypothetical protein EUROIMAGE1875335
208822_s_at	
218160 at .	gb:NN 014222.1 /DEF-Homo sapiens NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 8 (19kD, FGIV) (NDUFAB), mRNA. /FEA-mRNA /GEN=NDUFAB /PROD=NADH dehydrogenase (ubiquinone) 1 alphasubcomplex, 8 (19kD, PGIV) /DB_XREF=gi:7657368 — /UG-HS.31547 NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 8 (19kD, PGIV) /FL=gb:BC001016.1 gb:AF044953.1 db:NN 014222.1
221728_x_at	Consensus includes gb:AA628440 /FEA=EST /DB_XREF=gi:2540827 /DB_XREF=est:af26f02.s1 /CLONE=IMAGE:1032795 /UG=Hs.83623 nuclear receptor subfamily 1, group I, member 3

	1 TENY TOPN-1 VCF
	F-Human Chediak-Higashi syndrome protein short isoform (LYSI) mkNA, compiete cus. igashi syndrome protein short isoform /DB_XREF=gi:2654473 /UG=Hs.36508 Chediak-Higa
210943_s_at	
	09.1 /DEF=Homo sapiens, likely nomolog or yeast wilt, compounding or
	FLUZU4/9, clone MGC:1038, mRNA, complete cds. /FEA=mRNA /PROD=11kely homolog of yeast Nhp2, component of theHACA shoRNP;
	hypothetical protein FL/20479 /DB_XREF=g1:12652540 /OG=RS:25590 increcial protein remarks
209104_s_at	
	Gb:NM_019048.1 /DEF-Homo sapiens hypothetical protein (FLJ20/52), mKNA. /FRA-mKNA /CEN-FLD20/52 /FRA-mCNA /CEN-FLD20/52
217987 at	付
-	mit, ATPase, b (Psmcb), manh.
	, macropain) 26S
201699_at	(prosome, macropain) 26S subunit, ATFase, 6 /FL=gb:BC005390.1 gb:D/82/0.1 gb:AFU0b340.1 gb:Macropain) 26S subunit, ATFase, 6 /FL=gb:BC005390.1 gb:D/82/0.1 gb:AFU0b340.1 gb:Macropain) 26S subunit, ATFASE, 6 /FL=gb:BC005390.1 gb:D/82/0.1 gb:Macropain) 26S subunit, ATFASE, 6 /FL=gb:BC005390.1 gb:D/82/0.1 gb:AFU0b340.1 gb:D/82/0.1 gb:AFU0b340.1 gb:D/82/0.1 gb
	gb:NW_016451.1 / DEF=Homo sapiens coatomer protein complex, subunit beta (COFB), mana. / Fea.mana. / Sex-Corp.
201359 at	/PROD=coatomer protein complex, smount Deca / DE_Antr=gir//Joseph / Prod=
25-00-00-00-00-00-00-00-00-00-00-00-00-00	urinic
210027_s_at	/DB_XRRF=gi:178742 /UG=Hs.73722 APEX nuclease (multitunctional DNA repair enzyme) /ru-gu:bco4:0:1 ga:motes from
	4_000714.2 /DEF=Homo
	tein, transcript variant PBR, mRNA. /FEA=mRNA /GEN=BZRP /PROD=peripheral benzodiazapine receptor /DB_XREF=gi:6382068
202096_s_at	/UG-Hs.202 benzodiazapine receptor (peripheral) /fil-guibrovillor; garandovarapine receptor (peripheral).
70.00	
774740_5_ac	the many / 22 miles PROD-similar to the content of the content o
	C. elegens hypothetical 55.2 kD protein F16A11.2 /DB_XREF=gi:7657014 /UG=Hs.10729 hypothetical procein
200042_at	ch. NM 025126.1 / DEFE-Homo sapiens hypothetical protein FLJ21786 (FLJ21786), mRNA. /FRA=mRNA /GEN=FLJ21786
219035_s_at	
	gb:NM_004640.1 /DEF=Homo sapiens HLA-B associated transcript-1 (Dessie), inva. / Fear-inva. / Fe
200041 s at	transcript-1 /DB XREF=g1:4758111 /UG=Hs.55296 HLA-B associated transcript-1 /FL=gb:BC004350.1 gb:MacMullandur
112	Consensus includes gb:X03348.1 /DEF=Human mRNA for beta-glucocorticoid receptor (clone UBIO). /FEA-man. C. member 1
216321_s_at	/PROD=beta-glucocorticoid receptor / DB AKEE=gl:31001 / OG-AS:/3/12 interest receptor
	DEFENDING SEPTEMS DAY SCHOOL OF THE PRODE LEUKOCYTE-SPECI
214574_x_at	Hs.88411
	branched chain transacylase (£2 component of branched chain mens / FFR) franched franched
	yrup urine disease)
•	/DB_XREF=g1:4503264 /UG=Hs.139410 dihydrolipoamide branched chain transacylase (E2 component of branched chain
205370_x_at	Keto acid dehydrogenase complex; maple sylup uline disease; / / / / / / / / / / / / / / / / / / /
213872 at	Consensus includes gb:BE465032 /FEA=EST /DB_XREF=g1:9510807 /DB_XREF=est:nV/og09.x1 /LLOAR-IRAGE-21:950-

	gb:NM_000404.1 /DEF=Homo sapiens galactosidase, beta 1 (GLB1), mRNA. /FEA=mRNA /GEN=GLB1 /PROD=galactosidase, and galactosidase, beta 1 (FLE-ch.NM 000404.1 db:M27507.1 gb:M22590.1 gb:M34423.1
201576_s_at	Pred::10834965 /UG-Hs.79222 galactosidase, Deca
	change factor beta /DB_XREF=gi:4505572 /UG=Hs.172813 PAK-interacting exchange ractor
202548_s_at	/FL=gb:D63476.1 gb:NM 003899.1
	PROD=rab6 GTPses activating protein (GAP andcentrosome-associated)
204028_s_at	activating protein (dar and centiosome moscomestianists) / CLONE=NT2RP4001328
	P4504
202435_s_at	3688.1
	sapiens nypounetical process from 1.00-Hs.5080 hypothetical n FLJ20484 /DB_XREF=gi:8923447 /UG=Hs.5080 hypothetical
217980_s_at	1040.1 gb:NM_017840.1 gb:AF183428.1
,	gb:NM_001316.1 /DEF=Homo sapiens chromosome segregation 1 (yeast homolog)-like /DB_XREF=gi:4503072 /UG=Hs.90073 chromosome segregation 1 (yeast homolog)-like /DB_XREF=gi:4503072 /UG=Hs.90073 chromosome segregation 1
201112_s_at	
	JG=Hs.55498 geranylge
+6 2 66606	/PROD=gerany_gerany_upprospuece_s_record
202344_5_at	ch-in 024079. I (DEF-Homo saplens hypothetical protein MGC2840 similar to a purality glucosylinamisters (DR YREF-GI13129069
	FEA-mRNA /GEN-MGC2840 /PROD-hypothetical protein MGC2840 similar to aputative glucosyltransferase /FL-gb:BC001133.1 gb:NM_024079.1
203545_at	/UG=Hs.155356 hypothetical protein Mcc2040 Similar to a partial cds. /FEA-mRNA /GEN=KIAA0084 /DB XREF=gi:577298
24.07.46	Consensus includes gb:D42043.1 / DEF-number news -cz
777040-05	Consensus includes qb: AA887480 /FEA=EST /DB_XREF=gi:3003168 /DB_XREF=est:0]54812.51 /CLUNE-IRAGE:100011
212591_at	/UG=Hs.17428 RBP1-like protein
7000	
218005_at	Consensus includes gb: BE670915 /FEA=EST /DB_XREF=gi:10031456 /DB_XREF=est:7643a09.X1 /CLUNE=LEAGE:3203203
201165_s_at	/UG=Hs.153834 pumilio (Drosophila) homolog 1 /FL=gD:AF3153511 gD:AR/CLONE=IMAGE:2264634
212239 at	Consensus includes go:Albanist / Fear-Est / Discounty subunit, polypeptide 1 (p85 alpha) //UG=Hs.6241 phosphoinositide=3-kinase, regulatory subunit, polypeptide 1 (p85 alpha)
40 2 300700	Consensus includes gb:BF303597 /FEA=EST /DB_XKEr=g1:11204210 /DB_AXALT-CC.512000000000000000000000000000000000000
204300_3_at	Consensus includes gb:AI091079 /FEA=EST /DB_XREF=gi:3430138 /DB_XREF=est:qab3eUb.si /CLUME=imAUE:1030230
214853_s_at	/UG-Hs.81972 SHC (Src homology 2 domain-containing) Lemisioning Free-mRNA /PROD-estrogen sulfotransferase
210580 x at	gb:L252/5.1 /DEF=Human estroyen Surror mansferase family, cytosolic, 1A, phenol-preferring, member 3 /FL=gb:L224/3.1. //DB XREF=g1:463124 /UG=Hs.274614 sulfotransferase family, cytosolic, 1A, phenol-preferring, member 3 /FL=gb:L224/3.1. //DB XREF=g1:463124 /UG=Hs.274614 sulfotransferase family, cytosolic, 1A, phenol-preferring, member 3 /FL=gb:L224/3.1.
	s includes gb:AK
) 150 c at	
	F=Homo sapiens, Similar to putative methyltransferase, putative methyltransferase /DB_XREF=gi:12804482 /UG=Hs.
209265_s_at	/FL=gb:BC003031.1 gb:BC001650.1 gb:NM_019852.1

•	
	The same one 113 2 (Intra-Horno saniens vay 3 oncodene (VAV3), mRNA. /FEA=mRNA /GEN-VAV3 /PROD=vay 3 oncogene
	JD XREF=q1:7262390
218807_at	/UG=Hs.267659 vav 3 oncogene /FL=gb:AF067817.1 gb:AF118887.1 gb:NM_U05113.2
	gb:AF033026.1 /DEF=Homo sapiens bitunctional ATF sullLY rasedamics /FEA=mRNA /PROD=bifunctional ATP sulfurylaseadenosine5-phosphosulfate kinase /DB_XREF=gi:3378100 /FEA=mRNA /PROD=bifunctional ATP sulfurylaseadenosine5-phosphosulfate kinase / AF10527.1
209043_at	
	sapiens nypotnetica, pro- n FLJ22690 /DB_XREF=gi:13
1 0 00 00	gb:NM_012071.1 /DEF=Homo sapiens BUP protein (BUP), mRNA. /FEA=mkNA /GEN=BUP /FNOL-BUP /FNOL-BUP /FNOL-BUP /FNOL-BUP /FNOL-BUP /FL-db:AF078848.1 gb:NM_012071.1
218048_aL	sapiens mRNA for KIAA0593 protein, partial cds. /red-mkNA
00000)B_XREF=gi:3043709 /UG=HS.11861 005121.1
לחדאמם מר	gb:NM_005771.1 /DEF=Homo sapiens retinol dehydrogenase homolog (RDHL), mRNA. /FEA=mRNA /GEN=KUHL gb:NM_005771.1 /DEF=Homo sapiens retinol dehydrogenase homolog /FL=gb:AF067174.1
21.9799 s at	(care)
	eukaryotic translation initiation factor (effs) mean, compression factor /DB_XREF=gi:2558667 /UG=Hs.57783 eukaryotic translat
208688 x at	3, subunit 9 (eta, 116kD) /FL=gb:U78525.1
	no sapiens calpain 4, small
200001 at	subunit Subuni
	121
71010 740 0	/GEN=11M23 /FKUD=Lranslocase of inner mitochondrial membrane 23 (yeast) homolog /FL=gb:AF030162.1 gb:NM_U0b34/.1
	3_XREF=gi:545
201840_at	/UG=Hs.75512 neural precursor cell expressed, developmentally down-regulated 8 /FL-gp:D23004.1 gp.ma_corporation
38710_at	Cluster Incl. AL096/14: Home Sapiens mays, come control of the con
212287 at	Consensus includes gb:BF382924 /FEA=EST / DB_AKEF=g1:1130%313 / DB_AKEF=G1:1130%31 / DB_AKEF=G1:1130%313 / DB_AKEF=G1:1130%31 / DB_AKEF=G1:
	F=gi:4884378
203098_at	Y chromosome-like /FL=gb:AF081258.1 gb:AF081259.1 gb:NM 004824.1
218905_at	gb:NM 017864.1 /DEF=Homo sapiens Aypornetical process 1200510 /UG=Hs.279521 hypothetical protein FLJ20530 /FL=gq:NM 017864.1 /PROD=hypothetical protein FLJ20530 /DB XREF=gq:8923495 /UG=Hs.279521 hypothetical protein FLJ20530 /DB XREF=gq:8923495 /UG=Hs.279521 hypothetical protein FLJ20530 /DB XREF=gq:8923495 /UG=Hs.279521 hypothetical protein FLJ20530 /DB XREF=gq:NM 017864.1
218303 x at	gb:NM_016618.1 /DEF=Homo sapiens hypothetical protein (LOC3103), MACA 11 gb:AF217520.1 gb:NM_016618.1 tein /DE XREF=gi:7706155 /UG=Hs.5721 hypothetical protein /FL=gb:AF3645451 /CLONE=IMAGE:4650406
10000	Z
	ens hepatocellular carcinoma-associated antigen 60 (HCA00), MRVAT. 1 (HG-HS.30670 hepatocell) a-associated antigen 66 (DB XREF-gi:893731 (HG-HS.30670 hepatocell)
218715_at	FL=gb: AF244135.1 gb: AF116531.1
202184_s_at	gb:NM_018230.1 /DEF=Homo sapiens hypothetical protein FLMIU614 (FLMIU614), mux.: /.

	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
	in FLJ10814 /DB_XREF=gi:8922689 /UG=Hs.1245/ hypothetical protein flutuois /FlJ-
	<pre>gb:NM_003001.2 /DEF=Homo sapiens succinate dehydrogenase complex, subunit C, integral membrane procein, 1340 (SDNC), nuclear gene encoding mitochondrial protein, mRNA. /FEA=mRNA /GEN=SDHC /PROD=succinate dehydrogenase complex, nuclear gene encoding mitochondrial protein, mRNA. /FEA=mRNA /GEN=SDHC /PROD=succinate dehydrogenase protein, nuclear gene encoding mitochondrial protein, mRNA. /FEA=mRNA /GEN=SDHC /PROD=succinate dehydrogenase complex, subunit C, integral membrane protein,</pre>
202004_x_at	15kD /FL=gb:U57877.1 gb:NM_003001.2
. te a 250000	clone MGC:3182, mRNA, complete cds. /FEA=mkNA /PKOJ=UnkdOwn (7 Homo sabiens, clone MGC:3182, mRNA, complete cds /FL=gb:BCO
221044 s at	=qb:AB039903.1 gb:NM_021616.1
	Consensus includes gb:AL137335.1 /DEF=Homo sapiens mRNA; cDNA DKEZp434A179 (from clone DKFZp434A179); partial cds.
. 100000	
200000	aldo-keto reductase family 1, member Al (aldehyde reductase) (AKR1A1),
201900 s at	/GEN=AKRIA1 /FROD=aldo-keto reductase ramily 1, member Al (aldehyde reductase) /FL=gb:BC000670.1 gb:J04794.1 gb NM_006066.1 /UG=Hs.89529 aldo-keto reductase family 1, member Al (aldehyde reductase) /FL=gb:BC000670.1 gb:J04794.1 gb NM_006066.1
	gb:NM_015607.1 /DEF=Homo sapiens DKFZP547E1010 protein (DKFZP547E1010), mRNA. /FEA=mRNA /GEN=DKFZP547E1010 .
202560_6_at	_
	lactosic
201105_at	binding, soluble, 1 (galectin 1) /FI=gb:BC001693.1 gb:Ud45b-1 gb:NM_U02405-2.
204546 at	1
	Consensus includes gb:AI735692 /FEA=EST /DB_XREF=gi:5057216 /DB_XREF=est:at20b12.x1 /CLONE=IMAGE:235564/
214181_x_at	/UG=Hs.88411 lymphocyte antigen 117
210032 % 6#	gb:NM_014322.1 /DEF=Homo sapiens opsin 3 (encephalopsin) (UFN3), ukwa: /FEA-muka / GEN-CON / 2007 /
- V-250572	
205306_x_at	3-monooxygenase (Kynurenine 3-nydroxyrase) /in-gone occose-
202837 at	=gb:BC003553.1 gb:AB007447.1 gb:NM_006700.1
	gb:NM 016553.1 /DEF=Homo sapiens hypothetical protein (DKFZp547L134), mRNA. /FEA=mRNA /GEN=DXFZp54/L134 gb:NM 016553.1 /DEF=Homo sapiens hypothetical protein /FL=qb:AL162061.1 gb:NM_016553.1
202153_s.at	/PRODERSY INCIDED FOR THE PRINCIPLE OF T
209341_s_at	inhibitor of kappa light polypeptide gene enhancer in B-cells, kinase beta /FL=gb:AF031416.1 gb:AF08158.1
	Consensus includes gb:AI761110 /FEA=EST /DB_XREF=gl:5176/7/ /DB_XREF=est:W109MUZ.X1 /CLONE-IMAGE.230011 /CLONE-IMAGE.230011
212493_s_at	- 1
	1 /DEF=HOMO SADIENS B-CELL ASSOCIATED DIOCELL (NOW), MARKIS 150962.1 GB:NM_007273.1 GB:AF126=g1:6005853 /UG=Hs.7771 B-cell associated protein /FL=gb:AF150962.1 gb:NM_007273.1 gb:AF126
201600_at	
	gb:NW_004094.1 /DEF=Homo sapiens eukaryotic translation initiation ractor 2, subunit 1 (alpha, 35%) / DB_XREF=gh:4758255 / FEA=mRNA /GEN=EIF2S1 /PROD=eukaryotic translation initiation factor 2, subunit 1 (alpha, 35%) / DB_XREF=gh:4758255
ŕ	/UG=Hs.151777 eukaryotic translation initiation factor 2, subunit 1 (alpha, 35kD) /ku=gp:Bc002513.1 gp:002013.1
. 201144_s_at	gb:NM_004094.1
211985 s at	Consensus includes gb:Alb33/30 /FEA=E31 /Db_AABF-91.*/3/70 /DL_ABF.279009 matrix Gla protein
25-2-20-21-9	

pasup

	<pre>sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 0 (1972), ASHI) /DB_XREF=g1:4827 OD=NADH dehydrogenase (ubiquinone) 1 betasubcomplex, 8 (1982, ASHI) ogenase (ubiquinone) 1 beta subcomplex, 8 (1982, ASHI)</pre>
201226_at	b.AF077028.1 gb:NM_005004.1 gp:ALU80030.1
	apiens, /PROD=h: ld prot
208828_at	gb:aF226077.1 gb:NM_017443.1
204020_at	/UG=Hs.29117 purine-rich element binding protein A /FL=gb:M96684.1 gb:NM_005859.1
1040	gb:NM_021821.1 /DEF=Homo sapiens MDS023 protein (MDS023), mrwn. /FEA-mrwn /CIN-12021.1 // DEFE=G1:11141894 /UG=Hs.10724 MDS023 protein /FL=gb:AF182422.1 gb:NM_021821.1
75-75-077	Consensus includes gb:BF971923 /FEA=EST /DB_XREF=gi:12339138 /DB_XREF=est:602240326r1 /CLONE=IMAGE:432807.1 Consensus includes gb:BF971923 /FEA=EST /IDB_XREF=gi:12339138 /DB_XREF=est:602240136r1 /CLONE=IMAGE:432807.1
203514_at	DB_XREF=est:ab400
204334_at	/UG=Hs.21599 Kruppel-like factor 7 (ubiquitous) /FL=gb:AB015132.1 gb:NM_U03/U3.1 /UG=Hs.21599 Kruppel-like factor 7 (ubiquitous) /FL=gb:AB015132.1 gb:NM_U03/U3.1 /CLONE=IMAGE:3268504
203531 at	Consensus includes gb:BF435809 /FEA=ES1 /DB_AREF=91:11340124 /OE
ZOJJJI AC	pressor 1 (CHES1), mRNA.
205022_s_at	3=Hs.211773 checkpoint
	atype, 3
201532 at	55.1 gb:NM 002/88.1
	8051624 /UG=Hs.264981 2-50ligoadenylate
	/PROD=2-50ligoadenylate synthetase 2, isologin pri film p
204972_ac	/r_cr_grams_rai_r_grams_rai_r_s_s_s_s_s_s_s_s_s_s_s_s_s_s_s_s_s_s_
	VIG=Hs.78995 MADS box transcription
209200_at	eptide C (myocyte
	sapiens BTB (FOZ) domain Concaning 1 (2027)
217945_at	238.1 gb:AF355402.1
	XREF=g1:6912279 /UG=Hs.
201491_at	012111.1 gb:AF164791.1
	9078 immature
204868 at	/FILEGD:NM_001545.1
	F=Homo sapiens CGI-76 protein (LOC31632), mkWA: //Fra-mkWA //321-200331 gb:NR 01
217826_s_at	/DB_XREF=g1://u0311 / UG-RS:104-25 CG1 / DB_XREF=g1:11005587 / DB_XREF=est:AU144066 /CLONE=HEWBA1000798
212534_at	/UG-HS.285519 Homo sapiens cDNA FLJ11904 fis, clone HEMBB1000048
1 2 2 7 8 5 1 1 0	gb:BC000324.1 /DEF=Homo sapiens, Similar to granulin /FL=gb:BC000324.1 /orans=TMAGR:1019291 dranulin /DB XREF=gi:12653114 /UG=Hs.180577 granulin /FL=gb:BC000324.1
207779	4551075 /FEA=EST /DB
212188_at	/UG=Hs.109438 Homo sapiens clone 24775 mWNA sequence

	1
	gb:NM_004725.1 /DEF=Homo sapiens BUB3 (budding uninhibited by benzimidazoles 3, yeast) homolog (BUB3), mRNA. /FEA=mRNA /GEN=BUB3 /PROD=BUB3 (budding uninhibited by benzimidazoles 3, yeast) homolog /DB_XREF=gi:4757879 /UG=Hs.40323 BUB3 (budding uninhibited by benzimidazoles 3, yeast) homolog /FL=gb:BC005138.1 gb:AF047472.1 gb:AF053304.1 gb:AF081496.1
201458_s_at	gb:NM_004725.1 gb:NM_015339.1 /DEF=Homo sapiens activity-dependent neuroprotective protein (ADNP), mRNA. /FEA=mRNA /GEN=ADNP /PROD=activity-dependent neuroprotective protein /DB_XREF=gi:12229216 /UG=Hs.3657 activity-dependent neuroprotective pro-
201773_at	4 1 10
218751_s_at	Cal protein FL0110/1 / DB Aker=g1:09222031 / OG-ns:32235 hypometric From Sapiens flotillin 2 (FED712) nRNA. /FEA=mRNA /GEN=FLOT2 /PROD=flotillin 2 (FED712) hypometric flotillin 3 (FED712) hy
.201350_at	/UG=Hs.184488 flotillin / FL=gD:NNOU44/3.1 9D:NOO222.1 db:NM_004766.1 /DEF=Homo sapiens coatomer protein complex, subunit beta 2 (beta prime) (COPB2), mRNA. /FEA=mRNA
	/GEN=COPB2 /PROD=coatomer protein complex, subunit beta 2 (betaprime) /DB_XREF=gi:4758031 /UG=Hs.75724 coatomer protein complex,
201098_at	
	i hormone recep
703/32_at	sapiens pl
100000	l glycan, class C /DB_XREF=g1:4505/94 /UG=RS./5/90 phospharthythmostcar 5-10-70 j.NM 002642.1
202040_5_ac	gb:NM_002115.1 /DEF=Homo sapiens hexokinase 3 (white cell) (HK3), mRNA. /FEA=mRNA /GEN=HK3 /PROD=hexokinase 3 (white
205936_s_at	cell) /DB_XREF=g1:4504594 /UG=HS:15925) HeXOALMES (WILLO COLT) / MRNA. /FEA-MRNA /GEN-HDACI /PROD-histone deacetylase 1 https://www.nones.com/descentials/liber.com/nones.com/descentials/liber.com/nones.com/
201209_at	DB_XREF=gi:13128859 /UG=Hs.88556 histone deacetylase 1 /FL=gb:BC000301.1 gb:U500/3.1 gb:Nm_U04504:2 gb:D37207.5
1	Consensus includes gb:AI123426 /FEA=ESI / DB_ANEF=91:3335152 / DB_ANEF=51:3256.1 gb:AF180473.1 gb:NM_014515.1
21//98_ac	Consensus includes gb:BE963245 FEA=EST / DB_XREF=gi:11766663 / DB_XREF=est:601656874R1 / CLONE=IMAGE:3865699
209455_at	[VG=Hs.21229 I=Dox and WD-40 domain ID Fig. 1 Professions and State of the Community of the Professions and State of the Professions and Profession and Profession and Profession and Profession and Profession and Profes
	17/51.1 / DEF = NORD Septembrance of the control of
212397_at	CDNA DKFZD4341081Z (IIOM CIOLE DA APTITUTE) CONTROL (PECI), MRNA. /FEA-mRNA /GEN-PECI CONTROL OF THE CONTROL OF
	<pre>pnoy1-CoA isomerase /DB_XREF=g1:5174624 /UG=Hs.15250 perox1somal D3:DZ-enoy1-CoA isomerase /DB_XREF=g1:5174624 /UG=Hs.15250 /UG=H</pre>
218025_s_at	Homo sapiens sialyltrans
	EA=mRNA
203217 s at	3 €
	EST /DB_XRRF=g1:4988533 /DB_XREF=est:we38g03.XI /CLONE-IERGE:25333 /DB_XREF=est:we38g03.XI /CLONE-IERGE:25333
212812_at	domo saptens cura: 1027-113. (PEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (PPPIR8), mRNA. (FEA-mRNA gb:NM_002713.1 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Homo sapiens phosphatase 1, regulatory (inhibitor) subunit 8 (DEF-Hom
207830 s at	/GEN=PPPIRB /PROD=protein phospharase 1, regulatory (inhibitor) subunit 8 /FIL=gb:NM_002713.1 gb:U14575.1
	hase, X-linked (PRKX), mRNA. /FEA-mKNA /GEN-FRXA /FNON-FLOCKING.
204061_at	

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74 4 757510	Consensus includes gb:AV713720 /FEA=EST /DB_XREF=gi:10795237 /DB_XREF=est:AV713720 /CLONE=DCBBJA12 /UG=Hs.306434 Homo
	1 ~ ~
Z00000_dt	gp.NM_10183847.1 pp.NM_10183847.1 DEF=Homo sapiens hypothetical protein FLJ11296 (FLJ11296), mRNA. /FEA=mRNA /GEN=FLJ11296 PRODE PROTEIN PROTEIN
2188U5_ac	gb:NM_015640.1 /DEF=Homo sapens and the first tensor of the first tensor for the first form of the first tensor of the first form of the first tensor of the first form of the
217725_x_at	Dinding Protein / DB_XKKE=gi:/Octo23 /Oc-ms.103530 FAI-1 maker_binding protein / D SENSEARS / FRODEseryl-tRNA synthetase (SARS), mRNA. / FEA=mRNA / GEN=SARS / FRODEseryl-tRNA synthetase (SARS), mRNA. / FEA=mRNA / GEN=SARS / FRODEseryl-tRNA synthetase (SARS), mRNA. / FEA=mRNA / GEN=SARS / FRODEseryl-tRNA synthetase (SARS), mRNA. / FEA=mRNA / GEN=SARS / FRODEseryl-tRNA synthetase (SARS)
200802_at	Obsersus includes gb: Al633709 /FEA=EST /DB_XREF=EST: th71f03.x1 /CLONE=IMAGE:2124125 /UG-Hs.30174
213203_at	noismail nuclear activating complex, polypeptide 3, 1000 mana. /FEA=mana /GEN=SPOP /PROD=speckle-type POZ pro-
204640_s_at	
202324 s at	:NM_022735 1
, , , , , , , , , , , , , , , , , , , ,	EF=Human platelet act
Z11001_X_HL	sanions Nathanibi milinone oxidorednorase 39kDa sublimit mRNA, nuclear gene encoding
	/ DEF = ROUNG SAPIELS NADA - MANGALINGS CALCASS SANCE
208969 at	NADH denydrogenase (ubiquinone) i aipna subcomplex, 3 (39kD) /Fi=gb:AF050641.1 gb:NM_005002.1
202227_s_at	protein /FL=gb:AF016270.1 gb:NM_006696.1
7 - 18 B B 12C	Consensus includes gb:AW057781 /FEA=EST /DB_XREF=gi:5933420 /DB_XREF=est:wx03f09.x1 /CLONE=IMAGE:2542601 /UG=RS.29797 Fibosomal protein L10
1001174	
202545_at	9 /UG=Hs.155342 protein kinase C, delte /FL=gb:Lu/sou.i gb:Lu/sou.i gb:Lu/sou.
202214 s at	gb:NM_003588.1 /DEF=Homo sapiens cullin 4B (CUL4B), mRNA. /FEA=mRNA /GEN=CUL4B /FROD=CULLIN 4B /DB_AREF=91:132/0400 //OG=Hs.155976 cullin 4B /FL=gb:NM_003588.1 gb:AB014595.1
218366 x at	o sapiens hypothetical ein FLJ20859 /DB_XREF=
	gb:NM_017829.1 /DEF=Homo sapiens hypothetical protein FLJ20454 (FLJ20454), mRNA. /FEA=mRNA /GEN=FLJ20454 /PROD=hypothetical protein FLJ20454 /DB_XREF=gi:8923424 /UG=Hs.26890 hypothetical protein FLJ20454 /FL=gb:AF273271.1
218592_s_at	gD:NN U1023.1 gb:API645981. DEF=Homo sapiens cell division control protein 16 (CDC16) mRNA, complete cds. /FEA=mRNA /GEN=CDC16 pp:API645981. DEF=Homo sapiens cell division control protein 16 /DR XRFF=ri:553374 /UG=Hs.1592 CDC16 (cell division cycle 16, S. cerevisiae, ho-
209658_at	molog) /FL=gb:AF164598.1
	sapiens ES1 (zebrafish) protein, human homolog of (C110KF31), mkNA. /FEA=mkNA. otein, human homolog of /DB XREF=2011690 /UG=H221182423 ES1 (zebrafish) protei
202217_at	AA868332:a
54970_at	/gi=2963777 /ug=Hs.77978 /len=481

202441 at	799 /DB_XREF=est:AL568449 /CLONE=CS0DE001YC
. 211383_s_at	apiens mRNA; cDNA DKFZp434F2427 (from clone DKFZp434F2427); complete cds. hypothetical protein /DB_XREF=g1:6807664 /UG=Hs.27207 KIAA0982 protein /F
40 500 500	gb:NM_003365.1 /DEF=Homo sapiens ubiquinol-cytochrome c reductase core protein I (UQCRC1), mRNA. /FEA=mRNA /GEN=UQCRC1. /PROD=ubiquinol-cytochrome c reductase core protein I /DB_XREF=gi:4507840 /UG=Hs.119251 ubiquinol-cytochrome c reductase core protein I /FF=ch:1.16842.1 db:NM 003365.1 db:D26485.1
COSTOS	10 01
201827_at	/FIT=gb:106618.1 gb:NM_003077.1 gb:AF113019.1 gb:NM_006363.1 /DEF=Homo sapiens Sec23 (S. cerevisiae) homolog B (SEC23B), mRNA. /FEA=mRNA /GEN=SEC23B /PRGD=Sec23 (S.
201583_8_ac	
212696 s at	68633 /FEA=EST /DB_XREF=g1:12335848 /DB_XREF=est:602271068F1 /CLONE=IMAGE:4359209 protein 4
218229 s at	
210111 s at	apiens PNAS-138 mRNA, complete cds. /FEA-mRNA /Fs PNAS-138 mRNA, complete cds /FL=gb:AF277175.1
	sapiens non-metastatic cel tochondrial protein, mRNA. in /DB_XREF=gi:4505408 /1
100000000000000000000000000000000000000	(prosome, macropain) 26S subunit, non pain) 26S subunit, non-ATFase, 1 /DB_XR se, 1 /FL-cb:D44466.1 cb:NM_002807.1
201135 Sat	sapiens hypothetical protein FLJ20039 (FLJ20039), mRNA. /FEA=mRNA /GEN=FLJ20039 n FLJ20039 /DB XREF=di:8923045 /UG=Hs.267448 hypothetical protein FLJ20039 /FL=gbiNM_
70.000000000000000000000000000000000000	sapiens pro come (prosor subunit, no
75-2-75-75-75-75-75-75-75-75-75-75-75-75-75-	
220044 X at	db.ar. 1 / DEF=How cases ribeye mRNA, complete cds. /FEA=mRNA / PROD=ribeye / DB_XREF=gi:12034652 / UG=Hs.171391 gb.ar.22711.1 / DEF=How cases ribeye mRNA, complete cds. /FEA=mRNA / PROD=ribeye / DB_XREF=gi:12034652 / UG=Hs.171391 C-terminal binding protein 2 /FL=qb:AF222711.1 gb:NM_022802.1
	gb: BC004383.1 / DEF=Homo sapiens, Similar to polypyrimidine tract binding protein (heterogeneous nuclear ribonucleoprotein I), clone MGC:10830, mRNA, complete cds. /FEA=mRNA /PROD=Similar to polypyrimidine tract binding protein (heterogeneous nuclear ribonucleoprotein I) /DB_XREF=gi:13325139 /UG=Hs.172550 polypyrimidine tract binding protein
211271_x_at	-Homo prote
218998_at 201765 g.at	(FL=GD:AFI)341/.1 gD:NW_01.032.1 Consensus includes gb:AL523158 /FEA=EST /DB_XREF=gi:12786651 /DB_XREF=est:AL523158 /CLONE=CSODC001YM06 (3 prime)

	/UG=HS.119403 hexosaminidase A (alpha polypeptide) /FL=gb:NM_000520.2
	Consensus includes gb:AL080178.1 /DEF=Homo sapiens mRNA; cDNA DKFZp434K171 (from clone DKFZp434K171); partial cds. /FEA=mRNA /GEN=DKFZp434K171 /PROD=hypothetical protein /DB_XREF=gi:5262652 /UG=Hs.27194 DKFZP434K171 protein
204837_at	70kh molymentide (RNP
	gb:NM_003089.1 /DEF=Homo sapiens small nuclear illonucleoprotein 70kD polypepide(RNP antique) /DB_XREF=gi:4507118 /UG=Hs.174051 small nuclear ribonucleoprotein 70kD polypepide(RNP antique) /DB_XREF=gi:4507118 /UG=Hs.774051 small nuclear ribo
201221_s_at	polypeptide (RNP antigen) /rL=go:BCU00342.1 gp:mazasse.1 gs:mazasse.1 gs:mazasse.2 /FEA=mRNA /PROD=My027 protein /DB_XREF=g1:12001995 gb:AF061730.1 /DEF=Romo sapiens clone 016b03 My027 protein mRNA, complete cds. /FEA=mRNA /PROD=My027 protein /DB_XREF=g1:12001995
209092_s_at	/UG=Hs.279061 CGI-150 protein /FL=gb:AF061730.1 gb:AF151908.1 gb:NM 016080.1
202265_at	
203721 s at	gb:NM_016001.1 /DEF=Homo sapiens CGI-48 protein (LUCS1098), mkwa. /FEA-mkwa /sea-Eucsios / men-euchy / NGE-Hs. 6153 'CGI-48 protein /FL-gb:AF151806.1 gb:NM_016001.1
	gb:NM_006326.1 /DEF=Homo sapiens seven transmembrane domain protein (NIFIS14), mRNA. /FEA=mWNA /GEN=NIFIZ14 /FACU-SEVEN transmembrane domain protein /FL=gb:BC001118.1 gb:NM_006326.1
	041204 /FEA=EST /DB_XREF=gi:3280398 /DB_XREF=est:ov
213165_at	CDABPOU86 mRNA sequence Consensus includes ob:AL117643.1 / DEF=Homo sapiens mRNA; cDNA DKF2p434M245 (from clone DKF2p434M245). /FEA=mRNA / DB_XREF=q1:5912233
213198_at	/UG=Hs.5288 Homo sapiens mRNA; cDNA DKFZp434MZ45 (from clone DKFZp434M245)
212837 at	1464
22,00042	DEF-Homo sapiens adaptor-related protein complex 3, delta 1 subunit (AP3D1), mRNA. /FEA-mRNA /GEN-AF-3D1.
206592 s at	related protein complex 3, delta isubunit / DB_AKEF=g1:43013/8 / OG=63:70050 compres
	Consensus includes gb:NR_000532.1 /DEF=Homo sapiens propionyl Coenzyme A carboxylase, beta polypeptide (PCLB), nucleat yele encours, macconnection, mana. /FEA=CDS /GEN=FCCB /PROD=propionyl Coenzyme A carboxylase, betapolypeptide /DB_XREF=g1:4557043 /UG=Hs.63788
212694_s_at	propionyl Coenzyme A carboxylase, beta polypeptide /FL=gb:NM_000532.1
211759 x at	gp:bc003363.1 / Dar-inum Saprams/ Cross controllers/ (FL=db:BC005969.1
	Consensus includes gb:AL050205.1 / DEF-Homo sapiens mRNA, cDNA DKPZp586F1323 (from clone DKFZp586F1323). / FEA-mRNA / DE Arch-gi:rocarra
212714_at	/UG=HS.Zobij Homo Sairens navas, coar Dispose 2 / DB XREF = 1:13291862 / DB XREF = 1:002439888F1 /CLONE=IMAGE:4566380 /UG=Hs.84318 replication protein
201528_at	A1 (70kD) /FL=gb:M63488.1 gb:NM_002945.1
211795 s at	gb.aF198052.1 /DEF-Homo sapiens EVH1 domain binding protein mKWA, complete cos. /rika-bin. comp. protein (PYB-120130) /FL=gb:AF198052.1
219633 2 24	gb:NM_018394.1./DEF=Homo sapiens hypothetical protein FLJ11342), mRNA. /FEA=mRNA /GEN=FLJ1342 /FKOD=Lypothetical protein FLJ11342 /FL=gb:NM_018394.1
200800 c	I-4 0
	0.1 /DEF=Homo sapiens cDNA: FLJ23067 fis, clone LNG04993. /FEA=mRNA /DB_XREF=g1:10433938 /UU=ES:1
214719_at	, 12
213655 at	monocxygenasetryptophan 5-monocxygenase activation protein, epsilon polypeptide
	Consensus includes gb:AL049397.1 /DEF=Homo sapiens mRNA; cDNA DKFZp586C1019 (from clone DKFZp586C1019): / EEA-maak / DEF=Homo sapiens mRNA; cDNA DKFZp586C1019 (from clone DKFZp586C1019): / EEA-maak / DEF=Homo sapiens
212371_at	Homo sapiens mRNA; cDNA DXPZp586C1019 (from clone DXFZp586C1019)
	P051151.1
210166 at	
217958_at	

.

pa306

	279901 PTD009	
219269 at	/GEN=FLJ21616 /PROD=hypothetical	
219574 at	gb:NM_017923.1 /DEF=Homo sapiens hypothetical protein FLJ20668 (FLJ20668), mRNA. /FEA-mRNA /GEN-FLJ20668 /FROD-hypothetical protein FLJ20668 /FL-gb:NM_017923.1 FR-170668 /DR XREF-c1:8923612 /UG-Hs.12920 hypothetical protein FLJ20668 /FL-gb:NM_017923.1	
210183 x at	cotein SDK3 mRNA, complet protein /FL=gb:AF112222.1	.*
211582 x at	gb:AF000424.1 /DEF-Homo sapiens LST1 mRNA, cLST1C splice variant, complete cds. /FEA-mRNA /GEN-LST1 /DB_XREF-gi:2145063 /UG-Hs.8841.1 Lymohocyte antiqen 117 /FL-gb:AF000424.1	
212557 at	no sapiens mRNA for KIAA0576 protein, partial sprotein	
213387 at	sapiens mRNA for Klaal240 protein, partial cds. /FEA-mRNA /GEN-Klaal240 /PROD-KLAAl240 px rotein	
218143 s at	carrier membrane protein 2 (SCAMP2), mRNA. /FEA=mRNA /GEN=5 8030 secretory carrier membrane protein 2 /FL=gb:BC001376.1	
211615 s at	/DEF-Human leucine-rich protein mRNA, complete 177109 /FL-gb:M92439.1	
213359_at	i:1384833 /1 protein.1,	
217842 at .	n (LOC51631), mRNA. /FEA-mRNA /GEN-LOC51631 /FROD-CGI-74 protein /DB_XREF=94: b:NR 016019.1	, . , . , .
207170 s at	gb:NM_015416.1 /DEE=Homo sapiens DKFZP586A011 protein (DKFZP586A011), mRNA. /FEA=mRNA /GEN=DKFZP586A011 /PROD=DKFZP586A011 protein /DB XREE=qi:7661659 /UG-Hs.75884 DKFZP586A011 protein /FI=gb:NM_015416.1	: : : : : : : : : : : : : : : : : : :
214352 s at	EF=g1:11947594 /DB_XREF=est:602136427F1 /CLONE=IMAG	
	gb:NM_012199.1 /DEE=Homo sapiens eukaryotic translation initiation factor 2C, 1 (EIP2C1), mRNA. /FEM=mRNA /GEN=EIF2C1 /PROD=eukaryotic translation initiation factor 2C, 1 /PE_gqh:AF093097.1	
218287_s_at	ntigen (CAMPATH-1)	e e
205545 x at	ng factor similar to dnaj (SPF31), Hsp40) homolog, subfamily C, membe	· · · · · · · · · · · · · · · · · · ·
213750_at	Consensus includes gb:AA928506 /FEA=EST /DB_XREF=g1:3076797 /DB_XREF=est:cml7g03.s1 /CLONE=IMAGE:1541332 /UG-HS.10762 EST8	
221970 s at	Consensus includes gb:AUI58148 /FEA-EST /DB_XREF=gi:11019669 /DB_XREF=est:AUI58148 /CLONE=PLACE1011221 /UG=Hs.3211U5 Homo sapiens CLNA: FLJ21737 fis, clone COLF3396	
218684 at		
200628 s at .	/GEN-WARS /PROD-transfer RNA-Trp sygb:NM_004184.2	
	gb:NM_002582.1 /DEF=Homo sapiens poly(A)-specific ribonuclease (deadenylation nuclease) (PARN), mRNA. /FEA=mRNA /GEN=PARN /FROD=poly(A)-specific ribonuclease (deadenylation nuclease) /DB_XREF=gi:4505610 /UG=Hs.43445 poly(A)-specific ribonuclease (deadenylation nuclease)	
203905_at	ncer	
204039_at 63009_at	lm 1	
211922_s_at	gb:AY028632.1 /DEF=Homo sapiens catalase (CAT) mRNA, complete cds. /FEA=CDS /GEN=CAT /PROD=catalase /DB_XREF=gi:13562131 /FL=gb:AY028632.1	
212861_at	Consensus includes gb:BF690150 /FEA=EST /DB_XREF=gi:11975558 /DB_XREF=est:602186478T1 /CLONE=IMAGE:4298635 /UG=Hs.19210 E\$TS:	

	is transcription factor of /ri-gb:04394:
ar	gb: BE222709
213123_at · °	associated protein 3
204049_s_at	gb:NM_01472.1.1 /DEF FORMS Septems ANTIANO Gene product /FL=gb:AB014580.1 gb:NM_014721.1. //DB_XREF=g1:7662247 /UG=HS:102417 /VG=HS:101040 Gene product /RIANO196 / MRNA. /FEA=mRNA /GEN=KIANO196 /PROD=KIANO196 gene product
	gb:NM_014846.1 /DEF=Homo sappens X.AAA1130 years product /FL=gb:D83780.1 gb:NM_014846.1 /ING-HS.8294 KIAA0196 gb:NAA0196
at	gb:Ali36943.1 / DEP-Homo sapiens mans; class class construction of the construction of
	gb:NM 024633.1 /DEF=HOmo Sapiens Aypornetical protein FLJ21276 /FL=gb:NM 024633.1 FEA=mRNA /GEN=ARHGEF3 /PROD=Rho guanine FLJ21276 /DB_XREF=gi:13375863 /UG=HS.41502 hypochetical protein FLJ21276 /DB_XREF=gi:13375863 /UG=HS.41502 hypochetide exchange factor (GEF) 3 (ARHGEF3), mRNA. /FEA=mRNA /GEN=ARHGEF3 /FI=nh.AP249744.1
	/DEF=Homo sapien hange factor (GEF
218501_at	
212833_at	
209623_at	Consensus includes gb:AW45454 / FEATERS / COMPILE SP:AP310971.1 gb:AP301000.1 gb:NM 022132.2 Coenzyme A carboxylase 2 (beta) / FILESP:AB06049.1 gb:AP310971.1 gb:AP301000.1 gb:NM 022132.2 Coenzyme A carboxylase 2 (beta) / FILESP:AB06049.1 gb:AP310971.1 gb:AP301000.1 gb:NM 022132.2
 14 a 0,000 c	olog, S
219966 x at	gb:NW 017869:1 /DEF=Homo sapiens BANP homolog, SMAR1 homolog /FL=gb:NW 017869.1 DB XREF=gi:8923506 /UG=Hs:194637 BANP homolog, SMAR1 homolog /FL=gb:NW 017869.1
213275_x_at	THE IMAGE: 324203
210231_x_at	gb:D45198.1 /DEF-Human man a control of translocation (myeloid leukemia-associated) /FL-gb:143129.1 /UG-Hs:1663699 /UG-Hs:150275 alpha /DB_XREF-gi:912971 /UG-Hs:1663699 /UG-Hs:150275 alpha /DB_XREF-gi:912971 /UG-Hs:1663699 /UG-Hs:150275 alpha /DE_XREF-gi:9120271 /UG-Hs:1663699 /UG-Hs:150275 alpha /DE_XREF-gi:9120271 /UG-Hs:1663699 /UG-Hs:150275 alpha /DE_XREF-gi:9120271 /UG-Hs:1663699 /UG-Hs:150271 /UG-Hs:1663699 /UG-Hs:166369 /
212474 at	accembly 1-like, clone MGC:2171, mRNA, complete cds. /
25-27-27-27-27-27-27-27-27-27-27-27-27-27-	SF=Homo sapiens, Similar
208717_at	5.1 0 /PPA=EST /DB XREF=q1:4371526 /DB_XREF=est:tm39e01.x1 /CLONE=IMAGE:2160504 /UG=Hs.192789 EST\$
217527_s_at	Consensus includes gb:A44/0300 / Language Companination Warning ENTRY H.saplens to ALUG HUMAN ALU SUBFANILY SP SUDFACE CONTAMINATION WARNING FR/22625), mRNA. / FEA-mRNA / GEN-FL/22625 / PROD-hypothetical protein
220495 s_at	gb:NM 024715.1 / DEFENCED Septems 1376016 / UG=Hs.10634 hypothetical protein FLJ22625 / FL=gb:NM 024715.1 representational protein MGC:8454, mRNA, complete FLJ22625 / DB XRRF=gi:13376016 / UG=Hs.10634 hypothetical protein (transformer 2 Drosophila homolog) 10, clone MGC:8454, mRNA, complete FLJ22625 / DB XRRF=gi:13376016 / UG=Hs.10634 hypothetical protein FLJ22625 / Drosophila homolog) 10, clone MGC:8454, mRNA, complete
-	gb: BC000451.1 /DEF-Homo sapiens, spiroing ractor, arginineserine-rich(transformer 2 Drosophila homolog) 10 /DB_XKEF=91:12033302 /O-1204593.1 cds. /PEA=mRNA /PROD=splicing factor, arginineserine-rich(transformer 2 Drosophila homolog) 1 gb:u61267.1 gb:u61267.
200892_s_at	OD=fer-1 (C 51.1
201798_s_at	gb:NM_013421.1 / Der Endem Sages 70G-HS. 234680 fer-1 (C.elegans)-11ke 3 (myoferiin) / FL-gb:Artion 2007 1
202529 at	gb:MM_U01/00.1. DEFENDED STATES DEFENDED STATES DEFENDED DEFENDED DEFENDED STATES DEFENDED
	ated) /DB_XREF=gi:13097182 /UG=HS.100353 LECOM
208897_s_at	box polypeptine to the result of the provisional membrane protein 3 (35kD, Zellweger syndrome), clone not result of the protein 3 (35kD, gb:BC005375.1 /DEF=Homo saplens, peroxisomal membrane protein 3 (35kD, gb:BC005375.1 /DEF=Homo saplens, peroxisomal membrane protein 3 (35kD,
210296 s at .	PROD=peroxisomal membrane procein 3 (33KD), deilwey-c3 Zellweger syndrome) /FL=gb:BC006-Hs.241507 dibosomal protein Zellweger syndrome) /FL=gb:BC006-Hs.241507 dibosomal protein Zellweger syndrome) /FL=gb:BC006-Hs.241507 dibosomal protein
212460 at	Consensus includes gb:BE/38425 /FEA=ES1 /DD_Aum = S1.00 S6
7	

	ing enzyme UnpES (UNP) mRNA, complete cds. /FEA=CDS /GEN=UNP /FRUD=UnpES /D
211800_s_at	U ubiquitin specific processes (8.2 /DEF=Homo sapiens ATP-bincub-family D, member 3 /DB_XRE
202850_at	gb:H81182.1 Consensus includes gb:B6493972 /FEA=EST /DB_XREF=g1:13455486 /DB_XREF=est:602542252F1 /CLONE=IMAGE:4673316 /UG=Hs,16492 DKF2P564G2022 pro-
212202_s_at	ans lens epithelium-derived growth factor mRNA,
209337_at	factor /DB XREF=gi:3283351 /UG-Hs.82110 PC4 and SFRS1 interacting protein 1 /FL=gD:NN_021144.1 gi:Arcosydor. PGA=HRNA /PROD=TRAF family member-gb:NN_04180.1 /DEF=Homo sapiens TRAF family member-associated NFKB activator (PRNR), mRNA /FRA=mRNA /
207616_s_at	associated NFKB activator / DB_XREF=g1:4759249 / UG=HS.146847 TRAF Tamily member associated NFRA CEN=GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA. / FEA=mRNA GEN=GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA. / FEA=mRNA GEN=GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA. / FEA=mRNA GEN=GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA. / FEA=mRNA GEN=GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA. / FEA=mRNA GEN=GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA. / FEA=mRNA GEN=GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA. / FEA=mRNA GEN=GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA. / FEA=mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA. / FEA=mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA. / FEA=mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor complex member 1 (GOSH1), mRNA GOSH / PROD=g01gi SNAP receptor c
204630_s_at	member 1 /DB_XREF=gi:4758455 /UG=HS.8868 golgi SNAP receptor complex member 1 /FD=go.AFV/3520.1 9D:AFT_CT=0.1 SPECTOR OF a gene encoding two
	Consensus includes giverious of the familiar of the factor of the familiar of
	oraco de la complete cds. / PEN-mRNA sapiens, carnitineacylcarnitine translocase, clone MGC:1207, mRNA, complete cds. / PEN-mRNA (PROD-carnitineacylcarnitine translocase / DB_XREF-gi:12804552 / UG-Hs.13845 solute carrier family 25 (carnitineacylcarnitine translocase / DB_XREF-gi:12804552 / UG-Hs.13845 solute carrier family 25 (carnitineacylcarnitine)
203658_at	member 20 /FL=gb:BC001689.1 gb:NM_U0UJ8/.2 Poncensus includes db:BP690062 /FEA=EST /DB_XREF=g1:11975470 /DB_XREF=est:602186366T1 /CLONE=IMAGE:4298440 /UG=Hs.172550 pplypyrimidine
212015_x_at	tract binding protein (heterogeneous nuclear ribonucleoprotein I)
75 2 501000	Consensus includes gb:BG403671 /FEA=EST /DB_XKEF=g1:1329/119 /DB_XKEF=est:0024130311 /CEGGG GB:002011 /FEA=EST /DB_XKEF=g1:1329/119 /DB_XKEF=G1:0024130311 /CEGGG GB:002011 /FEA=EST /DB_XKEF=G1:1329/119 /DB_XKEF=G1:0024130311 /CEGGG GB:002011 /CEGG GB:002011 /CEGGG GB:002011 /CE
2021212 - S-01	Construction of the control of the c
2027.4	ens mRNA for pUb-R5, complete cds. /FEA-mRNA /GEN-hpUb-R5 /FROD=p
210460_s_at	ione male-greenific lethal-3 (Drosophila) -like 1 (MSL3L1), mRNA. /FEA-mRNA
207551_s_at	OB XREF=gi: 5803103 UG=HS: 88764 male-specific lethal-3 Urosoph
201104 x at	gb:NM_015383.1 /DEF=Homo sapiens hypothetical protein /FL=gb:NM_015383.1 DB_XREF=gi:7657016 /UG=Hs.218329 hypothetical protein /FL=gb:NM_015383.1
200812_at	viens chaperonin containing TCP1, subunit 7 (etc.) (CC17), mana. / ran-mana. / car. (CC17), mana. / car. (CC17), mana. / car. (CC17), mana. / car. (CC17), cubunit 7 (ct.) (EL=gb.AR0082). (ER=gi:5453606 / TCP1, subunit 7 (ct.) (ct.) (EL=gb.AR0082). (CC27), ct. (CC17), ct
203494 s at	gb:NM_014679.1
202127 at	Consensus includes gb: AB011108.1 /DEF=Homo sapiens mRNA for KIAA0536 protein, partial Cds. /FAM-man /Gan-Al-Andolo / John Jan 198891 serinethreonine-protein kinase PRP4 homolog /FL=gb:U481356.1 gb. 741620 / 103-Hs 15-485 huntingtin interact-
212532 g at	Consensus includes gb:AM873564 /FEA=EST /DB_XREF=gi:8007617 /DB_XREF=est:nob_CIU.XI /CLUCKE-IRAGE:JUXIJ) / CEA-AM873564 /FEA=EST /DB_XREF=gi:8007617 /DB_XREF=est:nob_CIU.XI /CLUCKE-IRAGE:JUXIJ) / LEA-AM873564 /LEA-AM87364 /LEA-AM8736
	gb:NM_012459.1 /DEF=Homo sapiens translocase of inner mitochondrial membrane 8 (Yeast) nomolog B (limanol), mars. Franslocase of inner mitochondrial membrane 8 (Yeast) homolog B /DE_XREF=g1:6912711 /UG=Hs.279915 translocase of inner mitochondrial membrane 8 (Yeast) homolog B /DE_XREF=g1:6912711 /UG=Hs.279915 translocase of inner mitochondrial membrane 8 (Yeast) homolog B /DE_XREF=g1:6912711 /UG=Hs.279915 translocase of inner mitochondrial membrane 8 (Yeast) homolog B /DE_XREF=g1:6912711 /UG=Hs.279915 translocase of inner mitochondrial
218357_s_at	
208847_s_at	gb:M29872.1 /DEF=Human alcohol dehydrogenase class III (ADHs) mkNA, complete cds. /FbA-mkNA /GEA-ALACA /GEA-TOOTON
207730_x_at	piens hypothetical protein FLJ20/00 (FLJ20/00), mans. /FAA-mans. /SAA-LLCCC hypothetical protein FLJ20/00 /FL=gb:NM 01/9932222 hypothetical protein FLJ20/00 /FL=gb:NM 01/99322 /PROD-KIAAO
212896_at	Consensus includes gb:D29641.2 / DRF-Homo saptens mRNA for KlAAUU32 protein, partial cus. / Landers db: 278608 KIAAU052 protein bb_XREF-gi:6633994 /UG-HS:278608 KIAAU052 protein protein
203156_at	anchor protein 11 /Fi=gb:AF16555.1 gb:NM_016248.1.
220066_at	gb:NM_022162.1 /DEF=Homo sapiens NOD2 protein (NOD2), mRNA. /FEA=mRNA /GEN=NOD2 /FROD=NOD2 protein /UB_AKEF=91:1137711 / OFF

	protein /FL=gb:AF178930.1 gb:NM_022162.1
218583_s_at	95:NM_020640.1 /DEF=Homo sapiens RF42 homolog (RP42), mRNA. /FEA=mRNA /GEN=RP42 /PROD=RF42 nomolog /DD_ALLE-51.101000000000000000000000000000000000
221689 s at	gb:AB035745.1 /DEF-Homo sapiens mRNA for DScK3D, Complete Cus. /Englands Cus. /En
	se.1 precursor /DB_XREF=gi:4557236 /UG=Ht
205412_at	acetyl-Coenzyme A acetyltransferase I (acetoacetyl controlly) mRNA. FFRA-mRNA gbing-to-coenzyme A acetyl-Coenzyme A man acetyl-Coenz
202542_s_at	member 1 (endothelialmonocyte-activating) /FL=gb:NM_004757.1 gb:U10117.1 member 1 (endothelialmonocyte-activating) /FL=gb:NM_004757.1 gb:U10117.1 member 1 (endothelialmonocyte-activating) /FL=gb:NM_004757.1 gb:U10117.1
202521 at .	gb:NM_006565.1 /DEF=Homo sapiens CCCT-Dinding factor (zinc finger protein) /FL=gb:NM_006565.1 gb:UZ5435.1 (zinc finger protein) /DB_XREF=gi:5729789 /UG-Hs.57419 CCCTC-binding factor (zinc finger protein) /DB_XREF=gi:5729789 /UG-Hs.57419 CCCTC-binding factor (zinc finger protein) /PROD=transcriptional intermediary
	gb:NM_015905.1 /DEF=Homo sapiens transcriptional intermediary factor 1 (TLT.), mrvn. /Fin-mrvn. /JDEF=Homo sapiens transcriptional intermediary factor 1 /Fin-gb:AF009353.1 gb:AF119042.1 gb:NM_003852.1 factor 1 alpha /DB_XREF=gi:7706233 /UG=Hs.183858 transcriptional intermediary factor 1 /Fin-gb:AF009353.1 gb:AF119042.1 gb:NM_003852.1
204391_x_at	gb:NW_015905.1
214733 s at	CONSCIENCE STATEMENT OF THE TABLE OF THE TAB
	(Asp-Glu-Ala-AspHis) box binding protein 1 /FI
217864_s_at	gb: AF167160.1 gb:NM_016166.1
212904 at	Consensus includes gb:Abb33011.1 /bs:-nound segizems maken consensus includes gb:Abb3301 /UG-Hs.198891 serinethreonine-
	Consensus includes gb:NA156948 /FEA=EST /DB_XREF=g1:1728563 /DB_XREF=EST:Z113102:51 /CLUML-LINGEST CONSENSUS Includes gb:NA156948 /FEA=EST /DB_XREF=g1:1728563 /DB_XREF=G1:1728561 /DB_XREF=G1:1728563 /DB_X
202120 ac	gb:NM_014302.1 /DEF=Homo sapiens Sec61 gamma (SEC61G), mRNA. /FEA-mRNA /GEN=SEC61G /PROD=Sec61 gamma /DB_AKEr=g1:/03/343 /GG=12:/03/343
203484_at	Gamma /FL=gb:R641841 g5:NM CD4302.1 Gamma /FL=gb:R641841 g5:NM CD4302.1
203345_s_at	Consensus includes gp:alloous /rea-bai / be.alloous / be.alloous / rea-bai / be.alloous / rea-bai / rea-ba
100000	Consensus includes gb:AI478147 /FEA=EST /DB_XREF=G1:43/13/3 /UB_AREF=ESC:LMD4LOCIAL /CONSENSUS MENNA /FEA=ERNA
213230_dc	The submit of th
	SGN=GFEEZ / PROD=general transcription factor IIE, polypeptide 2(beta-subunit, 34kD) / DB_Akbr=gi: 304L); / CON-CONTROL CONTROL CONT
202680_at	beta subunit,
218117_at	ens ring-box 1
218768_at	iens nuclear pore complex protein (Nofil), mass. 5204 nuclear pore complex protein $7E_{\rm c}$ -gb:NML020401
202271 at	Consensus includes gb:AB007952.1 / DEF=Homo sapiens mana lot Alanasas. Free=sp:3413925 / UG=Hs.64691 KIAA0483 protein /FL=gb:NM_015176.1
218543 s at	GD.NM. 022750.1 / DEF-Homo sapiens hypothetical protein FLJ22693 (FLJ22693), mkNA. / FEA-MANA / OLL-102750.1 FEA-FMRNA / GEN-GABBRI FLJ22693 / PE-90:AL136766
	<pre>gb:NM_001470.1 /DEF=Homb sapiens gamma-aminobutyric acid (GABA) B receptor, 1 (whereal, 10835014 /UG=Hs.167017 gamma-aminobutyric acid (GABA) B receptor 1, isoform a precursor /DB_XREF=gi:10835014 /UG=Hs.167017 gamma-aminobutyric acid (GABA) B receptor 1, isoform a precursor /DB_XREF=gi:10835014 /UG=Hs.167017 gamma-aminobutyric acid (GABA).</pre>
203146_s_at	receptor, 1 /FL=gb:NM_001470.1 gb:AF301005.1 gb:AF099148.1 mRNA /FEA=mRNA /GEN=APMCP1 /PROD=APMCP1 protein /DB XREF=gi:10864014
21,8140_x_at	gb:NW 021203.1 /DEF=Edomo sapiens Arger protein /Filegb:NW 021203.1 gb:AFILE Common of Common
40420 at	Cluster Incl. AB015718:Homo sapiens lok mkwa lor procesu Asmass, Compress / 18n=4221

	Consensus includes gb:AI928526./FEA=EST /DB_XREF=gi:5664490 /DB_XREF=est:wp59806.x1 /CLONE=IMAGE:2466034 /UG=HS.258730 hemp=regulated ini-
48580 at	
202416 at	gb:NM_003315.1 /DEF-Homo sapiens tetratricopeptide repeat domain 2 (17c2), manner 7 /FL-gb:Ud651.1 gb:NM_003315.1 2 /DB_XREF-gi:4507712 /UG-HS.5542 DnaJ (Hsp40) homolog, subfamily C, member 7 /FL-gb:Ud651.1 gb:NM_003315.1
211727 s at	ns, COX11 (yeast) homolog, cytochrome c oxidase ass homolog, cytochrome c oxidaseassembly protein (DB
	gb:NM_015049.1 / DEF-Homo sapiens amyotrophic lateral sclerosis 2 (juvenile) chromosome region, candidate 3 /DB_XREF=gi:13027379 /UG=HS.15/4248 amyotrophic (768N=ALSCRA) YROba-myotrophic lateral sclerosis 2 (juvenile) chromosome region, candidate 3 /DB_XREF=gi:13027379 /UG=HS.15/4248 amyotrophic (768N=ALSCRA) YROba-myotrophic lateral sclerosis 2 (juvenile) chromosome region, candidate 3 /Fleegb:RB038951.1 gb:NM_015049.1
202125_s_at 209451_at	Tarters Streets 2 105F=Human TRAF-interacting protein I-TRAF mRNA, complete cds. /FEA=mRNA /PROD=I-TRAF /DB_XREF=g1:151801 / VG=HS.I4004 / ASSECTION OF ASSECTION
209440 at.	ns, phosphoribosyl pyrophosphate Synthetese 1, trune morrisosyl pyrophosphate synthetase 1 /FL=gb:BC0016 phate synthetase 1 /DB_XREF=g1:12804406 /UG=Hs.56 phosphoribosyl pyrophosphate synthetase 1 /FEA=mRNA /GEN=SAS /
1000	<pre>gb.NX_018946.2 /DEF=Homo sapiens N-acetylneuraminic acid phosphate synthass; static acid phosphate synthase; stalic acid PROD=N-acetylneuraminic acid phosphate synthase /DB_XREF=gi:12056472 /UG=Hs.274424 N-acetylneuraminic acid phosphate synthase; stalic acid synthase /Phosphate 18946.2 qb:BC000008.1 gb:AP257466.1</pre>
200947_s_at	e 1 (GLUD1), mRNA. /FEA=mRNA (GEN=GLUD1 /FKU)=GLUCGGLUCASCUASC e 1 /FI=GR-2:013.48 L1 gb:M3715.41 gb:M3.0867.1 gb:NM_005271.1
100000	(S. cercvisiae) related yene ramary, among (S. cercvisiae) related B /DB_XREF=gi:5454045 /UG=Hs.7239 SEC24 (S. cercvisiae) related
205323 S.at	F-Homo sapiens metal-regulatory transcription factor 1 (MTF1), mRNA. /FEM-mANA / GEN-ALL. //DB_XREF=gi:5174588 /UG=Hs.211581 metal-regulatory transcription factor 1 /FL=gb:NM_0059
212622_at	Consensus includes gb:N64760 /FEA=EST /DB_XREF=gi:1212589 /DB_XREF=est:yz30c06.s1 /CLONE=IMAGE:284554 /UG=Hs.174905 KIAA0033 protein
212033_at	Consensus includes gb:BP055107 /FEA=EST /DB_XREF=gi:10809003 /DB_XREF=est:7175a05.xl /CLONE=IMAGE:3392240 /UG=Hs.180789 Sib4 procein
, 218919_at	gb.NM 024699.1 /DEF=Homo sapiens hypothetical protein First No. (First No. 1024699.1 PLJ14007 /DB_XREF=gi:13375984 /UG=Hs.99519 hypothetical protein fines frankorinion factor, complete cds. /FEA=mRNA /GEN=HPFF
209271_at	nns BPTF mRNA for Dromodomain fro 1119st temperature of 12 feetal Alzheimer antigen /FL=gb:ABC transcription factor /DB_XFR=gi:GbT (NG=FR:ABC) (NG=FR:ABC) feetal Alzheimer antigen /FL=gb:ABC (NG=FR:ABC) feetal Alzheimer antigen /FL=gb:ABC) feetal Alzheimer antigen /FL=gb:ABC (NG=FR:ABC) feetal Alzheimer antigen /FL=gb:ABC) feetal Alzheimer an
	Consensus includes gB:AL0317/8 / DEF=Human LAN Sequence Lical Consensus includes gB:AL0317/8 / DEF=Human LAN SEQUENCE LICAL CONSENSUS SERVED BZRP (peripheral benzddiazapine protein with 205 domain similar to part of Tight Junction Protein DNA sequence from clone 34B21 on chromosome 6pl2.1-21.1. Contains part of a gene for a PRRP (peripheral PRRP—RNNA_2 / DB_XREF=g1:4153588 / UG=HS.183056 Human DNA sequence from clone 34B21 on chromosome 6pl2.1-21.1. Contains part of a gene for a novel BZRP (peripheral hovel protein with ZUS domain similar to part of Tight Junction Protein 201 (TUPI) and UNCS Homologs, the gene for a novel BZRP (peripheral hovel protein with ZUS domain similar to part of Tight Junction Protein 201 (TUPI)
213322_at	benzodiazapine receptor
221895_at	Consensus includes gb:AW469184 /FEA=EST /DB_XREF=egi:7039290 /DB_XREF=est:hc78g04.x1 /CLONK=lRAGE:4030070 /UG=HS.14512 DIPB protein
217760_at	Consensus includes gb:AAL/0/80 /FEA=ESI /DE_ARGE-91:10:000 /FEA-91:10:000 /FEA-91:10:000 /FEA-91:000 /
201448_at	Consensus includes gb.AL046419 /FEA=ESI /DB_AKE=gi:3%34373 /D_AL046419 /FEA=ESI /DF.EA=ESI /DF.EA=E
218465_at	gb:NM_018126.1 /DEF=Homo sapiens hypothecical protein Fill0525 /FL=90:8000948.1 gb:NM_018126.1
212880_at	Consensus includes glos-Manians. / Jose-para de la consensus includes glos-Manians / Jose-para glos-para de la consensus includes glos-para de la consensus de
218142_s_at	016302.1 g
202078_at	gb:NM_00353.1 / DEF=AGMO Saprems COF9 (Constitutive photomorphogenic, Arabidopsis, homolog) subunit 3 /DB_XREF=gi:4502974 /UG=Hs.6076 COP9 (Constitutive photomorphogenic, Arabidopsis, homolog) subunit 3 /DB_XREF=gi:4502974 /UG=Hs.6076 COP9 (Constitutive photomorphogenic, Arabidopsis, homolog) subunit 3 /DB_XREF=gi:4502974 /UG=Hs.6076 COP9 (Constitutive photomorphogenic, Arabidopsis, homolog)

		T
	ens honothetical protein FLJ20189 (FLJ20189	
219069 at	gb:NM 01/104.1 / DEF-ROMD Septem 32052 hypothetical protein FLJ20189 /FL-gb:NM 017704.1 FROD=CGI-101 protein / DB XREF=gl 7705603 FLJ20189 / DB XREF=gl 7705603	33
	gb:NM 016041.1 /DEF=Homo sapiens CGI-101 protein (LOC31003), makes. / LDA-makes. /	,
218333_at	// Ud-mars. 2001.71 - 02 - 02 - 02 - 02 - 02 - 02 - 03 - 03	_
213082_s_at	DB XREF=gi:4008516 /UG=Hs.90078 nucleotide-sugar transporter similar to C. elegans sqy-/ DB XREF=gi:4008516 /UG=Hs.90078 nucleotide-sugar transporter similar to C. elegans sqy-/	1223 /
221452 s_at	gb:NM 0309b9.1 / DEFEROND Saprens 1250m.	
	Consensus includes gb:BF740152 /FEA-EST /DB_XREF=gi:12066828 /DB_XREF=est:7113908.x1 /CLONE=IMAGE:3564495 /00=n8.121303 mRNA. /FEA-mRNA /GEN	/GEN=SRM160
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201225_s_at	protein (plenty of prolines 101-like) /FL-gb:AF048977.1 gb:NM 005839.1 protein (plenty of prolines 101-like) /FL-gb:AF048977.1 gb:NM 005839.1	
219392 x at .	eat	shock pro-
	g	10.1
219212_at	gb:AF143723.1 gb:AF143723.1 gb:AF143723.1	:
217907 at	T=gb:AL136633.1 gb:BC001623.1 gb:AF161556.1 g	
		200
200727_s_at	(actin-related process 4, years) summary for A+U-rich element RNA binding factor, complete cds. /FEA-mRNA /FROD-A+U-rich element RNA binding factor / recomplete cds. /FEA-mRNA /FROD-A+U-rich element RNA binding factor / recomplete cds. /FEA-mRNA /FROD-A+U-rich element RNA binding factor / recomplete cds. /FEA-mRNA /FROD-A+U-rich element RNA binding factor / recomplete cds. /FEA-mRNA /FROD-A+U-rich element RNA binding factor / recomplete cds. /FEA-mRNA /FROD-A+U-rich element RNA /FR	filtering
209068_at	factor /DB_XREF=g1:3218539 /UG=Hs.170311 heterogeneous nuclear ribonucleoprotein D-11Ke /FL-9D:105005: 4 3:2000000000000000000000000000000000000	/len=684
46256 at	Cluster Incl. AA522670:ni39a05.s1 Homo sapiens CDNA, 3 end /clone=IMAGE-979184 /clone_end=3 /gy=An32301 /PROD=hypothetical protein	F
. Te 2	gb:NM 024836.1 /DEF=Homo sapiens Aypothetical protein FLJ22301 /FL=gb:NM_024836.1 FD MGC:1233) /DB XREF=gi:12655112	655112 /
2000017	db:BC001408.1 /DEF=Homo sapiens, clone MC:1233, mRNA, complete cds. /FEA=mrNA /FRUD-ULLIAMA (FLUD- db:BC001408.1 /DEF=Homo sapiens, clone MC:1233, mRNA, complete cds. /FEA=mrNA /FRUD-ULLIAMA (FLUD- db:BC001408.1 /DEF=Homo sapiens, clone MC:1233, mRNA, complete cds. /FEA=mrNA /FRUD-ULLIAMA (FLUD-ULLIAMA)	
208843_s_at	UG=HS.6880 DKFZP434D156 protein /FL=gD:BCU01406.1. UG=HS.6880 DKFZP434D156 protein /FL=gD:BCU01406.1. VFEA=mRNA /GEN=OAS1 / CANADAS / C	OAS1 /
	indud Septems 1, isoform E18 /DB_XREF=gi:8051620 /UG=Hs.82396 2,5-oligoadenylate Synthetase 1, isoform E18 /DB_XREF=gi:8051620 /UG=Hs.82396 2, isoform	
202869_at	/FL=gb:NM_016816.1 Concentis includes db:AL050378.1 /DEF=Homo sapiens mRNA; cDNA DKF2p58611420 (from clone DKF2p58611420); partial cds. /FrA=mRNA	
	-	for.
213546_at	DKRZDSTIRGINI PERSON / VOE KER-EST / DB_KREF-91:2265202 / DB_KREF-est:ng34d08.sl / CLONE=LMAGE:93000/ / VOE-ES-ES-ES-ES-ES-ES-ES-ES-ES-ES-ES-ES-ES	
214789_x_at		ctor,
213649 at	Horno sapiens	1S CDNA
	Consensus includes gb:AI972475 /FEA=EST / UB_AREF=91:3/03331 / UB_AREF=91:3/0331 / UB_AREF=91:3/031	
212251_at	gb:U68567.1 / DEF-Human lysosomal acid alpha-mannosidase mRNA, complete cds. /FEA-mRNA / PRUD-1950somal acid alpha-mannosidase mRNA, complete cds. /FEA-mRNA / PRUD-1950somal acid alpha-mannosidase mRNA, complete cds. /FEA-gb:NN 000528.1 gb:U65572.1 gb:U65572.1 gb:U65572.1 gb:U65572.1	
70,000	/DB_XREF=g1:1658373 /UG=Hs.2/9854 mannosidase, argue, case con ccoposes //Cr.ONP=TMAGE:3885630 /UG=Hs.108104 ubjquitin-db:U60266.1	i.
25-6-07-07-07	Consensus includes gb:BE964669 /FEA=EST /DB_XREF=G1:11/0820/ /DB_XREF=EST:002002002000 / Consensus includes gb:BE964669 /FEA=EST /DB_XREF=G1:110720	2090
200683_s_at	protein (LOC51340), mRNA. /FEA=mRNA /GEN=LOC51340	
219913_s_at	/UG-HS.268281 crooked neck protein (crn) /ru-guintzaurar garangen Frizaurar /GEN-FLJ20287 /PROD-hypothetical protein Prote	in
218104_at	PL.220287 / DB_XREF=g1:8923268 /UG=HS.26369 hypothetical protein FL.2028/ /FL=gD:NM_C1//*0:1	

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92	ludes gb: BE675139
212345_s_at DX	OKFZD586F2423
Co Co Co Co Co Co Co Co Co Co Co Co Co C	Consensus includes gribeivized frames for the following for form monnow (PROD=MondoA protein /DB_XREF=gi:7662347
	14938.1 /DEF=Homo sapiens KIAA0867 protein
202519_at //	IS CDC23 (cell division cycle 23, yeast, homolog) (CDC23), mRNA FEA-maka / GEN-CDC2 / FL-gb:AF053977. "In YRFF-ci:4757947 /UG-HS.153546 CDC23 (cell division cycle 23, yeast, homolog) /FL-gb:AF053977.
202892_at gl	b: AF191341.1 /************************************
213876 x at	Consensus includes gp:ANV05304 (Fig. 2) Consensus (Consensus Includes gp:ANV05304 (FROD-Aldehyde dehy-
	pb.NM_000382.1 /DEF=Homo sapiens aldenyde denydrogenase 3 family, member A2 /FL=gb:L47162.1 gb:U46689.1 gb:NM_U0U338.1.
(b)	gb.#F167438.1 /DEF=Homo sapiens androgen-regulated short-chain denydrogenaserreuctuse. / Jug-Hs.179817 CGI-82 protein /FL=gb:BC000112.1 /PR017 PR017 CGI-82 protein /FL=gb:BC000112.1 /PR017 PR017 CGI-82 protein /FL=gb:BC000112.1
217776_at	gb: AF151840.1 gb:NW_016026.1 gb: AF167438.1 / FEA-mRNA /
	onucleotiderransformylase /DB_XREF=gi:2317691 /UG=Hs.90280 5-aminoimidazore-w-v j.phvdrolase /FL=gb:U37436.1 gb:D82348.1 gb:D89976.1 gb:NM_004044.1
208758_at	ibonucleotide formyrightensiereseers 1970 DKF2P586C1620 protein
212749_s_at C	Consensus includes gb:AIO96477 /FERA-EST / DB XREF=91:3443971 / DB XREF=est:wt04405.x1 /CLONE=IMAGE:2506473 /UG=HS.250773 signal sequence
<u>د</u> ن	٠
200890_s_at	anslocon-associated r
	b.ar054589.1 /DEF=Homo sapiens nic procein isocorm in 1888 in Process in 1888 in Park in Park in 1888
	gb:NM_000923.1 /DEF=Homo saplens phosphodiesterase 4C, CAMP-specific (dunce(Drosophila)-homolog phosphodiesterase E1) /DB_XREP=gi 4505664
	PERALITY (SERVED DESCRIPTION OF SERVED OF SERV
206792 x at	ila)-homolog phosphodiesterase E1) /FL=gb:NM 000923.1
	pb:NM_016205.1 /DEF=Homo sapiens platelet derived growth factor C /FL=gb:AF091434.1 gb:AF244813.1 gb:AB035831.1
218718_at g	(splicing
	gb:NN 006924.1 /DEF=Homo sapiens splicing factor, arginine-rich 1(splicing factor 2, alternate splicing factor) /ub_Arar_glicing factor, /pron=splicing factor, arginine-serine-rich 1(splicing factor) /FL=qb:NG9040.1 gb:NM_006924.1
201742_x_at	UG-HS.73737 splicing factor, arginineserine-rich 1 (splicing ractor 2, arternace spreaments (GEN-MGC5585 /PROD-hypothetical protein MGC5585 /
	gb:NM_024057.1 /DEF=Homo saptens Aypothetical protein MGC5585 /FL=gb:BC000861.1 gb:NM_024057.1 DB_XREF=gi:13129027 /UG=HS.5152 hypothetical protein MGC5585 /FL=gb:BC000861.1 gb:NM_024057.1
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at	gb:NM_015960.1 /DEF=Homo sapiens CGI-32 protein (Notation) argini- UG-Hs.16606 CGI-32 protein /FL=gb:AP132966.1 gb:NM_01596.1 db:NM_01596.1 db
	bink_005626.1 /DBF=Homo sapiens Splicing ractor, arginineserine_rich 4 /FL=gb:BC002781.1 gb:L14076.1 go:RC00378.1 meserine_rich 4 /DB_XRBF=gi:5032088 /UG=HS:76122 splicing factor, arginineserine_rich 4 /DB_XRBF=gi:5032088 activating transcrip—neserine_rich 4 /DB_XRBF=gi:5032088 activating transcrip—
	Consensus includes gb:Al434345 /FEA=EST /DB_XREF=g1:4293929 /DB_AAM: Technology // Alfa Alfa Alfa Alfa Alfa Alfa Alfa Alfa
222103_at	tion factor 1 /r-gg:xm_court Concensus includes db:AW269335 /PEA=EST /DB_XREF=gi:6656365 /DB_XREF=est:xs47d05.x1 /CLONE=IMAGE:2/1/2/1/ /UG-ms:2/2/2/2/2/2/2/2/2/2/2/2/2/2/2/2/2/2/2/
204036_at	d G-protein-coupled receptor, 2 / rh=g0:0.0152:1 g5:00051188 / PROD=klaa-1so protein
218283_at	gbinM_U16305.1 /DEFinance Degree Fig. Dec. De
209654_at	gb: BC004902.1. / DEF=H0MO Saprems, C.

	1 (HCOROl) mRNA, complete cds. /rka=mkNA /skw=ncokkl /rkw=ccokkl /rkw=ncokkl /rkw=ncokkl /rkw=ncokkl /rkw=nkw /rkw=nkw 007074.1 gb:D44497.1
209083_at	/DB_XREF=gi:1002922 /UG=HS.109606 coronin, actin_Dinding process; au72d09.x1 /CLONE=IMAGE:2781809 /UG=HS.154095 zinc finger process for a process includes db:AM162015 /FEA=EST /DB_XREF=gi:6301048 /DB_XREF=est:au72d09.x1 /CLONE=IMAGE:2781809 /UG=HS.154095 zinc finger process
221873_at	
213573 at	Consensus includes gairwoolous (final final fina
1 n	gb.aF114784.1 /DEF=Homo sapiens methyl-CpG binding endonuclease (mm.) 4 /FL=gb:AF072250.1 gb:NM_003925.1 gb:AF114784.1 endonuclease /DB_XREF=gi:4539758 /UG=Hs.35947 methyl-CpG binding damain protein 4 /FL=gb:AF07147 /PROD=hypothetical protein MGC2747 /
ar	ob_Ander 91:12.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2
206513_at	/UG=HS.105115 absent in metanoma 2 / FE-90 contain 45 isoform (VPS45) mRNA, complete cds. /PEA-mRNA /GEN-VPS45 / PKULPVacuoiai pro-
209268_at	PR-3 /PROD=apoptosis
4	gD:NM_U10003.1 / LER-LIAMS SEPTEMBER
219329_s_at	PRODEZINC LINGEL Process
202051 s at	DISTRICT HOUSE / DISTRICT FIL-GD: AB007885.1 gb: NM. 005095.1 UCLONE-IMAGE: 3523665 /UG-HS.10842 RAN, member RAS UG-HS.150390 zinc finger protein 262 /FL-gb: Associated for the contract of t
	Consensus includes gb:BF112006 /FEA=EST /DB_XREF=g1:10941019 /DB_XREF
200749_at	oncogene family /
-	Sis-related Argential Property (FEED) 198143.1
213373_S_GL	formers includes db:D42084.1 /DEF=Human mRNA for KIAA0094 gene, partial cds. /rea-munn /GEN-Aimnova /
212673_at	KIAA0094 protein KIAA0094 protein
	gb:NM_005698.1 /DEF=Homo sapiens Secretory carrier membrane protein 3 /FL=gb:BC00161.2 gb:BC005135.1 gb:AFU05039.1 ne protein 3 /DB_XREF=gi:5032076 /UG=H8.200600 secretory carrier membrane protein 3 /FL=gb:BC00161.2 gb:BC005135.1 gb:AFU05039.1
201771_at	
	gb:U08032.1 /DEF=Human thermolabile (monoamine, m loim) phenolsulfotransferase (DB_XREF=gi:468256 /UG=Hs.274614 sulfotransferase family, cytosolic, lA, penolethermolabile (monoamine, M form) phenolsulfotransferase (DB_XREF=gi:468256 /UG=Hs.274614 sulfotransferase family, cytosolic, lA, penolethermolabile (monoamine, M form) phenolsulfotransferase (DB_XREF=gi:468256 /UG=Hs.274614 sulfotransferase family, cytosolic, lA,
209607_x_at	phenol-preferring, member 3 /FL=gb:NM 003166.1 gb:L19956.1 gb:U34193.1 gp:U34193.1 gb:U04034.1 phenol-preferring, member 3 /FL=gb:NM 003166.1 gb:L19956.1 gb:L1995
717317 S at	Consensus includes gb:ABU02391.2 /DE-name out. 2015.00 Consensus /DENT-XTAAAA83 /PROD-KIAAA483 protein /DB_XREF-gi:7662157 /
202272 s at	gb:NM_015176.1 /DEF=Homo sapiens KIAA0483 protein (KIAA0483), mkwa. /Fra-mkwa /sra-mkwa /crw-fsn1/DB XREF=q1:2145065 /UG-Hs.88411 UG-Hs.64691 KIAA0483 protein /FL=gb:NM_015176.1
16 % 003010	gb:AP000425.1 /DEF-Homo sapiens LST1 mRNA, cLST1A splice Variant, complete cus. /lin-marin 4 /DB XREF-q1:4507144 /UG-Hs.267812
27-07017	gb:NW 003794.1 /DEF=Homo sapiens sorting nexin 4 (SNX4), MRNA, /KRA=mrunA (GEN-SNA4 / ENCE-SCEENS)
205329_s_at	
218194_at	gD:NM_U15543.1 / DEF=noun Septems Inclease /FL=gb:AF151872.1 gb:AL110239.1 gb:NM_015543.1 DEFE SEPTEMS SEPTEM
220560 at	gb:NM_014144.1 /DEF=Homo Sapiens Smap Process (2007) / Described Smaps / FEA-man /GEN-ENPEP / FROD-arginyl aminopepti-
208270 s at	gb:NN_020216.2 /DEF=Homo sapiens arginy1 aminopepinase (aminopepidase (aminopepidase B) /FL=gb:NN_020216.2 dase (aminopepidase B) /DB_XREF=g1:13443030 /UG=Hs.283667 arginy1 aminopepidase (aminopepidase B) /DB_XREF=g1:13443030 /UG=Hs.283667 arginy1 research (100567397) /FEA=mRNA /GEN=LOC57397 /PROD=hypothetical
220175 s at	10a1 1907(
212429_s_at	Consensus includes gb:AM194657 /PEA=EST /DB_XKEF=g1:04/333/ /DB_XKEF=g1:04/33/ /DB_XKEF=g1:0

	FEAR-mRNA /PROD-iNMFF11hha /DB_XREF=gi:6526354 /UG=HS:174050 endothelial
100058 at	gb.AB002282.1 /DEF-Homo sapiens miNA ior immeration, compress and differentiation-related factor 1 /FL=gb.NM_003782.1
212535_at	Consensus includes gb:AA142929 /FEA=EST /DB_XREF=gi:1712307 /DB_XREF=est:2140g07.s1 /CLONE=IMAGE:504444 /UG=HB:288993 ESTS / COnsensus includes gb:AA142929 /FEA=mRNA /GEN=UBEZN /
•	gb.NM_003348.1 /DEF=Homo sapiens ubiquitin-conjugating enzyme EZN (nombio- pson=nbiquitin-conjugating enzyme EZN (bomologous toyeast UBC13) /DB_XREF=gi:4507792 /UG=Hs.75355 ubiquitin-conjugating enzyme EZN (nombio-
201524_x_at .	gous to yeast UBC13) /FL=gb: D83004.1 gb: BC000396.1 gb: DC003365.1 gb: DC003365.1 gb: NM_U03348.1.
201716 at	gb:NW 003099.1 /DEF=Homo saptens Soluting mexim 1. Spi-NF1065483.1 gb:NW 003099.1 Soluting nexim 1 /FL=gb:BC000357.1 gb:U53225.1 gb:NF1065483.1 gb:NF1065483.1 gb:NF1067851.1 gb:NF106785483.1 gb:NF1067854838483.1 gb:NF10678548384838484848484848484848484848484848
10000	
772037 S ac	333 /FEA-EST /DB_XREF-gi:5363390 /DB_XREF-est:Wn/9elu.xi /chown-limosi.zovov
213532_at	to ORF YOR126c S.cerevisiae trachea cellular apoptosis susceptibility protein (CSE1) mRNA, complete cds. /FEA-mRNA /GEN-CSE1
	protein /DB XREF=gi:3560554 /UG=Hs.90073 chromosome segregation 1 /Yeast name 159
210766_s_at	/FL=gb:AF0536401.1 Home sapiens capiens CDNA FLJ10756 fis, clone NT2RP3004572, highly similar to Home sapiens conjector of 1111-
209523_at	.1 gb:AF04
209276_s_at	glutaredoxin (thioltransferase) /FL=gb:BC005304.1 gb:AF162769.1 gb:DZ1238.1 glutaredoxin (thioltransferase) /FL=gb:BC005304.1 gb:AF162769.1 gb:AF163.1 /CLONE=IMAGE:754288 /UG=Hs.4113 S-adenosylhomocysteine
200848 at	٠,
	58.5
212406_S_at	/DEF-Human DNA sequence from clone RP5-863C/ on chromosome 2021.1.1.
212072_s_at	Kinase 2 alpha 1 polypeptide 12 2:1762327 /UG=Hs 239189 glutaminase C /DB_XREF=gi:7662327 /UG=Hs 239189 glutami-
	gintaminase (923), mai: < 1 ab: AF097493.1 ab: AF22
203159_at	ns endocrine regulator (HRIHPB2436), mRNA. /FEA=mRNA /GEN=HRIHFB24
203521_s_at	JOB KREF=g1:7657183 /UG-HS. 48433 endocrine regulator /FL=gD:Arianianianianianianianianianianianianiani
	ist /DB_XREF=gi:5803026 /UG=Hs.75470 hypothetical protein, expressed in Osteoblast /ill-grinse
204439_at	gb:NM_006820.1
218646 at	gb:NM_01/86/11 (DEF=ncm0 Saprems in formation protection FLJ20534 (FL=gb:NLJ36673.1 gb:NM_01/86/1.1 FLJ20534 (DB:NREF=gi:8923502 /UG=HS.44344 hypothetical protect FLJ20534 (FL=gb:NLJ20534 /UG=HS.44450 Sp3 transcription fac-
	Consensus includes gb:AUI45005 / FEA=ESI / LD AKEK-giracosto
213168_at	ton Physical NDEF-Homo sapiens, Similar to retinoblastoms binding protein 4, close MGC11139, many compression of PL and MGC1011.
210371_s_at	to retinoblastoma binding protein 4 (DB_XREF=gi:13111850 (UG=HS.15003 retinoblastoma premarkA (GEN=ILF2 /PROD=interleukin enhancer to retinoblastoma envisor interleukin enhancer binding factor 2, 45kD (ILF2), mRNA. /FEBN=RRNA /GEN=ILF2 /PROD=interleukin enhancer binding factor 2, 45kD (ILF2), mRNA. /FEBN=RRNA /GEN=ILF2 /PROD=interleukin enhancer binding factor 2, 45kD (ILF2), mRNA. /FEBN=RRNA /GEN=ILF2 /PROD=interleukin enhancer binding factor 2, 45kD (ILF2), mRNA. /FEBN=RRNA /GEN=ILF2 /PROD=interleukin enhancer
200052_s_at	gb:U10323.1 GENERATION Sapiens eukaryotic translation initiation factor 1A (EIFIA), mRNA. /FEA-mRNA /GEN-ELFIA / FRU-ELFIA / F
	tion initiation factor 1A /DB_XREF=gi:4503498 /UG=Hs.4310 eukaryotic translation initiation factor 1A /DB_XREF=gi:4503498 /UG=Hs.4310 eukaryotic
201019_s_at	gb:NML 001412.1
219892 at	family member 1 /DB_XREF=gi:13194198 /UG=Hs.133865 transmembrane 6 superiamily member 1 /Fb-Bire 1 /DB_XREF=gi:13194198 /UG=Hs.133865 transmembrane 6 superiamily member 1 /PEA-mRNA /GEN=LOC55858 /PROD-uncharacterized
	gb:NW 018475.1 /DEF-Homo sapiens uncharacterized Lypomeraterized hypothalamus protein HTMP /FL=gb:BC003545.1 gB:Arazuleo.1 hypothalamus protein HTMP /DB_XREF=gi:8923860 /UG=Hs.236510 uncharacterized hypothalamus protein HTMP /DB_XREF=gi:8923860 /UG=Hs.236510 uncharacterized hypothalamus protein HTMP /FL=gb:BC003545.1 gB:Arazuleo.1
218095 s at	gb:NN_018475.1 gb:Ar183409.1

	b 4 tribition of Comment County of the Principle of the County of the Co
	complex subunit 4 (LOC51138), mRNA. /FEA-mRNA /GEN=LOC51138 /FKOU=COF9 complex subunit 4 /FF-Th-Brond-102 - h. API-00757.1 gb:NM_016129.1
218042_at	28927 /DB_XREF=est:602324811F1 /
202172_at	161 /FL=gb:D28118.1 gb:NM_007146.1
	gb:NW_018108.1 /DEF=Homo sapiens nypocnetical protein FLJ10483 /FL=gb:NW_018108.1
218108_at	400
212539_at	FLJ22530 FLJ22530 TOPA-FRY (DR XREF=q1:6300659 / DB XREF=est:au68bll.x1 / CLONE=IMAGE:2781405 / UG=Hs.21739 Homo sapiens mRNA;
212635 at	Consensus includes grawing of Fig. 22. Consensus for the control of Consensus 1150695 (UG-Hs.7145 calpain 7 control of Co
	Consensus includes gb:BB349584 /FEA=EST /DB_XREF=G1:945143/ /DB_AREF=C30:00000000000000000000000000000000000
203356_at	ens UZ small nuclear ribonucleoprotein auxiliary factor, small subunic 2 (UZAFIRSZI), maren
	Schulzarists / Thomseleoprotein auxiliaryfactor, small subunit-related protein 2 /DB_XREF=gi:4827045 /UG=Hs.171909 UZ small nuclear
208174 x at	rkun-uz sugar increar factor, small subunit 2 /FL-gb: D49677.1 gb:NM_005089.1 ribonucleoprotein auxiliary factor, small subunit 2 /FL-gb:D49677.1 gb:NM_005089.1
	f10.x5 Homo sapiens cDNA, 3 end /clone=Image=301.03 /clone=
53912_at	1.10n=601 /10n=601 /10n one -Innon. 5.07286 /UG=Hs. 25431 KIAA1219 protein
221736_at	/GEN=UNPH4
	EF=gi:4504204 /UG=Hs.75782
1	/GEN=GTF3CZ / PROD=general transcription tactor files of FE-gb:D13636.1 gb:NM 001521.1
204366_s_ac	TION TOLOW T
212584_at	product product / PEA-mRNA / GEN-KIAA0648 protein, partial cds. / FEA-mRNA / GEN-KIAA0648 / PROD-KIAA0648 protein /
4 0 7 1 1 1 1	Consensus includes gb:Ab014340.1 / DEF-Edun Sayrum
21777	db:NM 018130.1 / DEF=Homo sapiens hypothetical protein FLJ10339 (FLJ1033) under / LLJ10330 1
219083_at	FLJ10539 /DB_XREF=g1:8922499 /UG=HS.93391 hypothetical protein FLJ10539 (from clone DKFZp586M2023); partial cds. /FEA-mRNA
015772 x at	Consensus includes gb:Aucovzzo: 1 / Dzr-nown organisms of 100-Hs. 247309 succinate-Coa ligase, GDP-torming, Deta subunit /GEN-DKFZp586MZ023 /PROD-hypothetical protein /DB_XREF-gi:4884469 /UG-Hs. 247309 succinate-Coa ligase, GDP-torming, Deta subunit
	Consensus includes 9b:AD001527 (DEF=Homo sapients DNA Iron cornensus 1-1-Cosamo 2000)
216194_s_at	/FEA-CDS_3 /DB_XREF=gi:1905899 /UG=BS.31003 CylcsAsteron mens. /FEA-mRNA /GEN-KIAA0480 /FROD=KIAA0480 gene product
204373_s_at	DB_XRRF=gi:7662155 /UG=Hs.92200 KIAAA0480 gene product /FL=gb:AB007949.1 gb:NM_014810.1 /DB_XRRF=gi:7662155 /UG=Hs.92200 KIAAA0480 gene product /FL=gb:AB07749.1 gb:NM_014810.1
1, 100000	Consensus includes gb:BF6/3888 /rEAmESI /DE_Aras-girilary / Jacks / Ja
	sapiens toll-like receptor 7 (LOC51284), mkNA.
220146_at	/DB_XREP=g1:7706092 /UG=HS:179152 toll-like receptor / /rb-gs.nrna. /FEA=mRNA /GEN=FLJ11127 /FROD=hypothetical protein
40,000	
100 M	131 /FEA=EST /DB_XREF=g1:11682455 /DB_XREF=est:nab19e04.x1 /CLONE=IMAGE:326638
213229_ac	CONSTRUCT TO THE TOTAL OF THE T
214356_s_at	Consensus includes plantification for the part of the property of the protection of the property of the protection of the property of the protection of the
213153 at	Consensus includes 92.505.25 XIAA1076 protein DB_XREF=gi:5689488 /UG=Hs.154525 XIAA1076 protein
100010	gb:AF267865.1 DEF=Homo sapiens DC41 mkNA, complete Cus: //Lineary Complete Cus: //File624.1 JPH=-01:AF267865.1 JPH=-01:AF267865
75-8-867017	
221522 at	/PROD=hypothetical

10,00

	protein /DB_XREF=g1:12053080 /UG=AS:39230 nome server TTR nolvoeptide 1 (alpha subunit, 56kD) (GTF2E1), mRNA. /FEA-mRNA
	SD-NM_005513.1 / DEF-Homo saplens general transcription factor 1.2, ports 5031726 / UG-HS.145381 general transcription factor (/ I/GEN-GTF2E1 /
	l transcription factor IIE, polypeptide L(alpha surmitt). Spride
205930_at	11s, polyperciae 1 (alpha subunit, 56kD) /FL=gb:NM_005513.1
210312 s at	gb:BC002640.1 /DEF=Homo Sapiens, Junia to doctor 24636 /FL=gb:BC002640.1 (cin /DB_XREF=gi:12803610 /UG=Hs.4187 hypothetical protein 24636 /FL=gb:BC002640.1
1 7	s1 /CLONE=IMAGE: 525848 / UG=RS. 2/3000 AND CLOSES FOR CONTROL OF STREET AND CONTROL OF
	ens mutl (E. coll) homolog 1 (colon cancer, mongolyposis type 2) /FL=gb:NM_000249.1 //UG=Hs.57301 mutl (E. coll) homolog 1 (colon cancer, nonpolyposis type 2) /FL=gb:NM_000249.1
202520_s_at	gb:U07418.1
221834_at	consensus includes grant of the state of the
210942_s.at	DB_XREF=gi:482724
202396_at	063 transcription
10.0000	gb:NW_016104.1 /DEF=Homo sapiens Fibula protein (Fibous);
	gb:NM_000016.1 /DEF-Homo sapiens acyl-Coenzyme A dehydrogenase, C-4 to C-12straight chain proenzyme /DB_XREF=gi:4547230
202502_at	/UG-Hs.79158 acyl-Coenzyme A dehydrogenase, C-4 to C-12 straight chain /rug-gb:000Next /CLONE-IMAGE:2508468 /UG-Hs.289088 heat shock 90kD
211968_s_at	Consensus includes gb:A1962333 /FEA=E31 /DE_ALLE_SILLOSSES / DE_ALLE_SILLOSSES / DE_ANOME / DECANOME / DECANOM
218027_at	gb:NM_014175.1 / DEF=Homo saptens Harling process proc
217792_at	gb:NM_014426.1 / DEF=Ramo Sapreis Soltwis Soltwis 1 Sp. NM_014426.1 sorting nexts / FEA=mRNA /GEN=DKFZP564D0372 / Sorting nexts / Tree=Eamen searies hypothetical protein DKFZP564D0372 (DKFZP564D0372), mRNA. /FEA=mRNA /GEN=DKFZP564D0372 /
221036_s_at	gb:NW U31301.1 / DEF = ROLL OF STREE = STREE = GI:13775223 /FILE = STREE = CONTROL OF STREE = CONTROL OF STREET OF S
	gb:U41514.1 / DEF=Numan UDF-Grand. DOF-Grand. DOF-Grand. DOF-HS.80120 UDF-N-acetyl-alpha-D-galactosamine:DOF-Grand. DolypeptideN-acetylgalactosaminisferase / DB_XREF=gi:1136284 / UG=HS.80120 UDF-N-acetylalactosaminisferase / DB_XREF=gi:1136284 / UG=HS.80120 UDF-N-acetylalactosaminyltransferase / DB_XREF=gi:1136284 / UG=HS.80120 UDF-N-acetyl-alpha-D-galactosamine:DOF-MS-ACETYL UDF-N-acetylalactosamine:DOF-MS-ACETYL UDF-N-acetylalactosamine:DOF-MS-ACETYL UDF-N-acetylalactosamine:DOF-MS-ACETYL UDF-N-acetylalactosamine:DOF-MS-ACETYL UDF-N-ACETYL UDF-N-AC
201/23_S_ac	Consensus includes gb:AIO93579 /FEA=EST /DB_XREF=gi:3432555 /DB_XREF=est:qDaJgV0.XI /CDCNL-includes gb:AIO93579 /FEA=EST /DB_XREF=gi:3432555 /DB_XREF=est:qDaJgV0.XI /CDCNL-includes gb:AIO93579 /FEA=EST /DB_XREF=est:q
202351_at	alpha V (vickonectin receptor, a.g.m. complete cds. /FEA-mRNA /PROD-calnexin /DB_XREF=g1:306480 /UG=Hs.135500 callexin /Inggreen.cds. /FEA-mRNA /PROD-calnexin /DB_XREF=g1:306480 /UG=Hs.135500 callexin /Inggreen.cds.
208853_s_at	gb: BC003552.1 gb: M94859.1 gb: M98452.1 gb: L10284.1 gb: L10284.1 gb: L10284.1 gb: M94859.1 gb: M94859.1 gb: M94859.1 gb: M94859.1 gb: M94859.1 gb: M94859.1 gb: M94869.1 gb: M94869.1 mailides db: AM014788 / PEA=EST / DB_XREF=g1: 5863545 / DB_XREF=est: UI-H-BIO-aae-h-10-0-UI. sl / CLONE=IMAGE: 2709354 / UG=Hs. 48802 Homo
213278_at	Sapiens clone 23632 mRNA sequence sapiens / DB XREF=q1:11018529 / DB_XREF=est:AU157008 /CLONE=PLACE1005711 /UG=HS.193725 proteasome (prosome,
203447_at	Consensus includes Baracia No. 18 S. F.Lgb:NM 005047.1 macropain) 26S subunit, non-Argass / FLL-gb:NM 005047.1
212459 x at	Consensus includes gb:Br393940 /FEA-E31 /JE2, ALL ST. 10587 /FEA-MRNA /GEN-FLJ10587 /PROD-hypothetical protein
218514 at	hypotnetical
212648_at	Consensus includes gb:ALU/323.1. / Dec-name september 2 Arabidopsis thaliana /DB_XREF=gi:5102732 /UG=HS.95665 Nypothetical process. General Consensus for the control of th
200708 at	gb:NN_002080.1 /DEF=Homo Sapiens Firemer Common PROD-aspartate aminotransferase 2 precursor /DE_ARGE=91.450400/10 PROD-aspartate aminotransferase 2) /FL=gb:BC000525.1 gb:NN_002080.1 Glutamic-oxaloacetic transaminase 2, mitochondrial (aspartate aminotransferase 2) /FL=gb:BC000525.1 gb:NN_002080.1 Glutamic-oxaloacetic transaminase 2, mitochondrial (aspartate aminotransferase 2) /FL=gb:BC000525.1 gb:NN_002080.1
20.004	

pa306

	ens poly(A)-binding protein, cycoplasmic 4 (inductible form) (inductible form) (inductible form) / DB_XREF=gi:6552335 /UG=Hs.169900 poly(A)-binding prote
201064_s_at	اھ
	gb:NM_014319.2 / DEF=Homo sapiens integral inner nuclear membrane protein /FL=gb:AF112299.2 gb:NM_014319.2
218604_at	86.1 /DEF-Homo sapiens cDNA FLJ14324 fis, clone PLACE4000100, highly similar to Homo sapiens hydr
214864_s_at	micha / DB_Arch: glitty-group for managers and analog (CDC16), mRNA. /FEA-mRNA /GEN=CDC16 /
	tens corto (cera monolog) /DB_XREF=g1:4502700 /UG=Hs.1592 CDC16 (cell division cycle 16, S. cerevisiae, homo.
202717_s_at	ens putative zinc finger protein (LOC51780), mRNA.
201643_x_at	
213803_at	lens hematopoletically expresse
204689_at	homeobox /DB_XREF=gi:10835016 /UG=Hs:118651 hematoportationally expressed independent respirators 2 /DB_XREF=gi:13376841 /UG=Hs/280776 tanky-
218228 S at	gb: AF
25-2-2019	sapiens CGI-95 protein mRNA, complete cds. /FEA=mRNA /PROD=CGI-95 protein /ub_Arcr=g1:4723030 /00-03:
202918_s_at	F=gi:6360061 /DB_XR
212263_at	
	Consensus includes gb:AU151793 /FEA=EST /DB_XREF=g1:11013314 /DB_XREF=esc:AU13173 /CLOUP-ALEACTOR / CONSENSUS INCLUDES gb:AU151793 /FEA=EST /DB_XREF=g1:11013314 /DB_XREF=esc:AU13173 /CLOUP-ALEACTOR / CONSENSUS INCLUDES gb:AU151793 /FEA=EST:AU13173 / CLOUP-ALEACTOR / CONSENSUS INCLUDES gb:AU151793 / FEA=EST:AU13174 / CLOUP-ALEACTOR / CONSENSUS INCLUDES gb:AU151793 / FEA=EST:AU13174 / CONSENSUS INCLUDES gb:AU13174 / CONSENSUS INCLUDES gb:AU1317
221825_at	FLOIL/19 ILS. TONE NEW TONE NEW MGC:12798, MRNA, COMPLETE Cds. /FEA-mRNA /PROD=Unknown (protein for MGC:12798) /DB_XREF=g1:13623468
211047_x_at	
209798_at	Locus (FL=gp:D0543.1 gp:NRT_0225.7. de RA70, complete cds. /FEA=mRNA /GEN=RA70 /PROD=RA70 /DB_XREF=gi:4062959 /UG=Hs.52644 SKAP55 homolo-db:AB014486.1 /DEF=Homo sapiens mRNA for RA70, complete cds. /FEA=mRNA /GEN=RA70 /PROD=RA70 /DB_XREF=gi:4062959 /UG=Hs.52644 SKAP55 homolo-db:AB014486.1 /DEF=Homo sapiens mRNA for RA70, complete cds. /FEA=mRNA /GEN=RA70 /PROD=RA70 /DB_XREF=gi:4062959 /UG=Hs.52644 SKAP55 homolo-
. 204361_s_at	gue /FL-gb: BC002893.1 gb: AE072166.1 gb: AE014486.1 gb: AF051323.1 gb: AF0513230.1 gb: AF051323.1 gb: AF051323
i d	Consensus includes gb:AI804118 /FKA=EST / DB_AKER = 91:0303030 / DB_AKER = 91:030300 / DB_AKER = 91:03030
218127_at	iens hypothetical protein
220355_s_at	mRNA. /FEA=mRNA /
	gb:NM_003850.1 /DEF=Homo sapiens succinate-CoA 11gase, ADF-1011111, Deca Succinate-CoA ligase, ADF-forming, beta subunit /FL=gb:NM_003850.1 ligase, ADF-forming, beta subunit /FL=gb:NM_003850.1
202930_s_at	gp:ABU33303.1 Consensus includes gb:AI801013 /PEA=EST /DB_XREF=gi:5366485 /DB_XREF=est:Wg15409.X1 /CLONE=IMAGE:2365169 /UG=Hs.211571 holocytochrome c
203745_at	Synthase (cytochrome c heme-lyase) /FL=gb:U36787.1 gb:NM_005333.1 synthase (cytochrome c heme-lyase) /FL=gb:NM_005333.1 ransla-
- C0300C	Consensus includes gb:All13320 / FEA=EST / DB_ANEF-91:333300 / SE_STONES 1 gb:U78311.1 gb:NM_003750.1 FEA=EST / DB / D
78-16007	EA=mRNA /GEN=FLJ20/30 05207.1 qb:NM_017945.1
218519_at	FLJ2017 (DB.XXEE-gg12556 / UG-HS.21460 INFOCHERICAL PLOCETH ILLEADY AS 3777 (CSTF3), IRWA. /FEA-IRWA / GEN-CSTF3
	gb:NM_001340:1 / DEFENCIO Saprens Creaves Comment 3, 77kD 77kD
203947_at	GENERAL OCASSOL 1 DEPEROND Sapiens hypothetical protein FL721901 (FL721901), mRNA. /FER-mRNA /GEN-FL721901 /FROD-hypothetical protein
219002_at	PLJ21901 /DB_XREF=g1:13375843 /UG=Hs.32546 hypothetical protein FLJ22344 (FLJ22344), mRNA /FEA=mRNA /GEN=FLJ22344 /PROD-hypothetical protein
220122_at	1_gb:NW_024717.1 3. camma (48kD) (ISGF3G), mRNA. /FEA=mRNA /GEN=ISGF3G
40.000	~1
203882 at	

	- 1.
212174_at	/PROD=s (18kD)
202567_at	gb:NM_0041/3.1./Dpr-name September 18kD) /DB_XREF=gi:4759159 /UG=Hs.15/5 small nuclear libouratectrons of polypeptide(18kD) /DB_XREF=gi:9910459 /UG=Hs.15627 Nit /FEL=gb:BC000457.1 gb:BC003150.1 gb:NM_004775.1 gb:U15009.1 /REL=gb:BC000457.1 gb:DC02150.1 gb:NM_004775.1 gb:U15027 Nit /FEL=gb:BC000457.1 gb:DC02150.1 gb:NM_004775.1 gb:U15027 Nit /FEL=gb:SD=Nit protein 2 /DB_XREF=gi:9910459 /UG=Hs.15627 Nit
218557_at	gb:NM_020202.1 /DEF=Homo Sapiens Mil process 1 20202.1 protein 2 /FL=gb:AF260334.1 gb:AF260334.1 gb:MG-020202.1 protein 2 /FL=gb:AF260334.1 gb:AF260334.1 gb:MG-020202.1 protein 2 /FL=gb:AF260334.1 gb:MG-020202.1 protein 2 /FL=gb:AF260334.1 gb:MG-020202.1 protein 2 /FL=gb:AF260334.1 gb:MG-020202.1
	3.1 / DEF-ENDANC SEPTEMBER SEPTEMBER SEPTEMBER CDNA: FL721862 fis, clone HEPU2121, Highly Samilar CONFEGURATION (NEFFECTION 30931 / UG-HS.188882 Homo sapiens CDNA: FL721862 fis, clone HEPU2121, HIGHLY SAMILAR CONFEGURATION (NEW-PORMIZ) / PRODECT
211976_at	ins cAMP responsive element binding protein-like 2 (CREBLZ), mana.
201990_s_at	sive element binding protein-line 2 / LL L L L L L L L L
208956_x_at	gb:062891.1 / DEF FRUMEN GEOXY TO THE PYTOPHOSD HAS FEED FILED: AD049113.1 gb:031930.1 gb:06281.1 gb:06281.1 gb:06281.1 gb:06281 FEED FIRE FEED FI
213939 s at	Consensus includes go: A10.1041 (Fig. 22) (Fig. 22) (Fig. 23) (Fig
201198 s at	Consensus includes gb:A1860431 /FEA=EST / DB_ARGE=g1:0.14 / DE ANN 002807.1 macropain) 26S subunit, non-ATPase, 1 /FE=gb:D44466.1 gb:NN 002807.1 macropain) 26S subunit, non-ATPase, 1 /FE=gb:D4466.1 gb:NN 002807.1 macropain) 26S subunit, non-ATPase, 1 /FE=grand of strains of transcription 5A (STAT5A), mRNA. /FEA=mRNA /GEN=STAT5A //FROD=signal
	gb:NM_003152.1 /DEF=Homo sapiens signal transcurer and activator of transcription of transcription of transcription of transcription of transcriptions /DB_XREF=gi:4507256 /UG=Hs.167503 signal transcribe and activator of transcriptions /DB_XREF=gi:4507256 /UG=Hs.167503 signal transducer and activator of transcriptions /DB_XREF=gi:4507256 /UG=Hs.167503 signal transducer and activator of transcriptions /DB_XREF=gi:4507256 /UG=Hs.167503 signal transducer and activator of transcription of transcriptions /DB_XREF=gi:4507256 /UG=Hs.167503 signal transducer and activator of transcriptions /DB_XREF=gi:4507256 /UG=Hs.167503 signal transducer and activator of transcriptions /DB_XREF=gi:4507256 /UG=Hs.167503 signal transcriptions /DB_XREF=gi:4507256 /UG=Hs.167
203010_at	-mRNA /6
218067_s_at	GD:NM_L10011.1 / District Control of the Control of
219077 s at	gb:NM_0163/3.1 / Def =name Sagrems "" CL=gb:AF211943.1 gb:NM_016373.1 gb:AF2Z/32/1.1 Gb:AFZZ/32/1.1 Gb:AFZZ/32/1 Gb:AFZZ/32/1 Gb:AFZZ/32/1 Gb:AFZZ/32/1 Gb:AFZZ/32/1 Gb:AFZZ/32/1 Gb:AFZZ/32/1 Gb:AFZZ/32/1 Gb:AFZZ/32/1
215424 s at	Consensus includes gb:AV689564 /FEA=EST /DB XREF=gi:10291427 /DB XREF=est:AV689564 /CLONE=GKCEAAU0 /UG=nS::35U0 37. CONSENSUS includes gb:AV689564 /FEA=EST /DB XREF=gi:10291427 /DB XREF=gi:10291427 /DB XREF=est:AV689564 /CLONE=GKCEAAU0 /UG=nS::35U0 37. CONSENSUS includes gb:AV689564 /FEA=EST /DB XREF=gi:10291427 /DB XREF=gi:10291427 /DB XREF=set:AV689564 /CLONE=GKCEAAU0 /UG=nS::35U0 37. CONSENSUS INCLUDES GRAND / GEN=MGC3062 /PROD=hypothetical protection MGC3062 /
219043 s.at	gb:NM_024065.1 /DEF=Homo sapiens Aypornetical Flores MGC3062 /FF=gb:AF267853.1 gb:CM01021:1 gb:NM_024065.1 pnding protein 1 / DB_XREF=gi:13129043 /UG=Hs.94576 hyporhetical protein 1, complete cds. /FEA=mRNA /GEN=XAB1 /FROD=XPA binding protein 1 /
209313_at	gb. AB044661.1 /DEF=Homo saptens AAb1 mww lot are controlled to protein /FL=gb.AB0446b1.1 Def=Home Saptens AB0
202944 at	gb:NW 000262.1 / DEFERONC Saprens N-acety-Manactorsaminidase, alpha- /FL-gb:BC000095.1 gb:Mb2/83.1 gb:Nb2.003.1 gb:Nb2.003.1 gb:Nb2.003.1 gb:Nb2.003.1 gb:Nb2.003.1 gb:Nb2.003.1 gb:Nb2.003.1 mgNb2.003.1 gb:Nb2.003.1 mgNb2.003.1 gb:Nb2.003.1 mgNb2.003.1 mgNb2.
1 0 0 0 0	gb. BC002719.1 / DEF=Homo sapiens, eukaryout character 3, subunit 1 (alpha, 35kD) / DB_XREF=g1:12803762 / UG=HS.17390, eukaryout character and attent
211656 x_at	gb:M3557.1 / DEF=Human MHC HLA-DQ beta mRNA, complete cds. /FEA=mRNA /GEN=HLA-DQB1 /DB_XREF=gi:6650833 /UG=Hs.83126 TATA box binding pro-
209358_at	gb:AF118094.1 /DEF=Homo Sapiens FRU2134 MRNA, Compresse 11, 12, 28th /FL=Gb:D63705.1 gb:NM_005643.1 gb:AF118094.1 /FEA-mRNA /GEN-DEGS tein (TBP) -associated factor, RNA polymerase 11, 12, 28th /FEL=Gb:D63 Discoppilla; lipid desaturase) (DEGS), mRNA, /FEA-mRNA /GEN-DEGS
	log prosoputa;
	E=g1:10284299 /DB_XREF=est:AV682436 /CLONE=GKBABEU8 /UG=BS:3040 Mumb Set:AFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF
. 213070_ac	port 1 / DEF Homo sapiens brain cellular apoptosis susceptibility protein (1921) 11ke 1 / DEF Homo sapiens susceptibility protein / DB_XREF=gi:3560556 / UG=Hs.90073 chromosome segregation 1 (yeast homolog) - 11ke Definity apoptosis susceptibility protein / DB_XREF=gi:3560556 / UG=Hs.90073 chromosome segregation 1 (yeast homolog) - 11ke Definity apoptosis susceptibility protein / DB_XREF=gi:3560556 / UG=Hs.90073 chromosome segregation 1 (yeast homolog) - 11ke DEF=Homology - 11ke DEF=Homolog
201111_at	/FL-gb:013286.1 gb:AF-053641:1 gb:NM 001316.1 /PEA-mRNA /FEA-MRNA /GEN-HEAB /PROD-ATPGTP-binding protein /DB_XREF-gi:5803028
204370_at	gb:\nm_U0651.1 / Lil

	ACUS TARREST STATE THAT THE TARREST STATE
202911_at	/DB_XREF=gi:4504190 /UG=Ms.3248 muts (E. coli) homolog 6 /FL=gb:U28946.1 gb:BC004246.1 gb:NM 000179.1 gb:U54777 2
204197_s_at	gb:NM_004350.1 /DEF=Homo sapiens runt-related transcription factor 3 (RUNX3), mRNA. /FEA=mRNA /GEN=RUNX3 /PROD=runt-related transcription factor 3 /Pri=ch-NM 004350 1
217809_at	gb:NM_014038.1 /DEF=Homo sapiens HSFC028 protein (HSFC028), mRNA. /FEA=mRNA /GEN=HSFC028 /PROD=HSFC028 protein /DE_XREF=gi:7661743 /UG=Hs.5216 HSFC028 protein /FL=gb:AP10323.1 db:BC003056.1 db:AP083246.1 db:AP083
201054_at	Consensus includes gb:BE966599 /FEA=EST /DB_XREF=gi:11772191 /DB_XREF=est:601660283R1 /CLONE=IMAGE:3906044 /UG=Hs.77492 heterogeneous nu- clear ribonucleoprotein AO /FL=gb:BC001008.1 gb:NM 006805.1 db:U73803.1
200994_at	Consensus includes gb:BG291787 /FEA=EST /DB_XREF=g1:13050002 /DB_XREF=est:602386007F1 /CLONE=IMAGE:4515240 /UG=Hs.5151 RAN binding protein / /FL=gb:AF098799.1 gb:NM_006391.1
201572_x_at	gb:NM_001921.1 /DEF=Homo sapiens dCMP deaminase (DCTD), mRNA. /FEA=mRNA /GEN=DCTD /PROD=dCMP deaminase /DB_XREF=gi:4503276 /UG=Hs.76894 dcMP deaminase /FL=gb:L12136.1 gb:NM_001921.1
	gb:NM_024573.1 /DEF=Homo sapiens hypothetical protein FLJ12910; mRNA. /FEA=mRNA /GEN=FLJ12910. /PROD=hypothetical protein FLJ12910;
218195_at .	OI .
209362_at	Apocaretical Process
203983 <u>a</u> t	
212074_at	4 (FEAREST /DB XREF=ci:10586110 /DB XREF=set:6015500/851 //r comp_reser.
204859_s_at	lens apoptot ivating fact 7.1 gb:NM 01
205707_at	gb:NM_014339.1 /DEF=Homo sapiens interleukin 17 receptor (1b17R), mENA. /FEA=mENA /GEN=1b17R /FROD=interleukin 17 receptor //Interpression 100 XREF=gi:7657229 /UG=HS.129751 interleukin 17 receptor /Pi=ch:n58917 1 ch:nm 014330 1
218716_x_at	gb:NW_012123.1 /DEF=Homo sapiens CGI-02 protein (CGI-02), nRNA, /FEA-mRNA /GEN=CGI-02 /PROD=CGI-02 protein /DB_XREF=gi:6912299 /UG=Hs.33979 CGI-02 protein /FL=gb:AF319422.1 gb:AF132937.1 qb:NM 012123.1
217940_s_at	Sign Sign
219802_at	GD:NM_024854.1.7DEF=Homo sapiens hypothetical protein FLJ22028 (FLJ22028), mRN. /FEA-mRNA /GEN=FLJ22028 /PROD=hypothetical protein FLJ22028 /FI=ch:NM 024864 1
209259_s_at	ns chromosome-associated polypeptide (HCAP) m EF=gi:3089367 /UG=Hs.24485 chondroitin sulfat
212037_at	Consensus includes gb:BF508848 /FEA=EST /DB_XREF=gi:11592146 /DB_XREF=est:UI-H-BI4-aor-e-06-0-UI.sl /CLONE=IMAGE:3085907 /UG=Hs.44499 pinin, desmosome associated protein
211063_s_at.	gb.BC006403.1 /DEF=Homo sapiens, NCK adaptor protein 1, clone MGC:12668, mRNA, complete cds. /FEA=mRNA /PROD=NCK adaptor protein 1 / DB_XREF=gi:13623576 /FL=gb:BC006403.1
203791_at	gb:NM_005509.2 / DEF=Homo sapiens Dmx-like 1 (DMXL1), mRNA. /FEA=mRNA /GEN-DMXL1 /PROD=Dmx-like 1 /DB_XREF=g1:9961348 /UG=Hs.1181042 Dmx-like
203405_at	ens Down s 4505022 /U
201604_s_at	ens myosin phosphatase, target subunit 1 (MYPT1), mRNA. /FEA-mRNA /GEN-MYP/ /UG-H8.16533 myosin phosphatase, target subunit 1 /F1=ch.ww 002480 1
218348_s_at	gb:NW_014153.1 /DEF=Homo sapiens HSFC055 protein (HSPC055), mRNA. /FEA-mRNA /GEN-HSPC055 /PROD-HSPC055 protein /DB_XREF-gi:7661761 //UG-HS.179898 HSPC055 protein /FL-gb:AF161540.1 gb:NM_014153.1
205659_at	gb:NM_014707.1 /DEF=Homo sapiens histone deacetylase 7B (HDAC7B-PENDING), mRNA. /FEA=mRNA /GEN=HDAC7B-PENDING /PROD-histone deacetylase 7B /FL=cp:AB018287.1 cp:NN 014707.1
202268_s_at	r protein-binding pro//DB_XREF=gi:4502168 /

	F LCCCCC
5	10.1 gb:U50939.1 gb:Na_u003003.1 sigma 2 subunit (AP352), mRNA. /FEA=mRN
o H	is adaptor-related process complex 3, a sabror-related protein complex 3, a 2subunit /DB_XREF=g1:5031580 /UG=Hs.154782 adaptor-related protein complex 3,
202399_s_at /	protein (LOC51325), mRNA. /FEA-mRNA /GEN-LOC51325 /PROD-mypounelical processing to the protein (LOC51325), mRNA. /FEA-mRNA /GEN-LOC51325 /PROD-mypounelical processing to the protein of the processing to the pro
218515_at /	/FL=gb:AF2U88b2.1
209180 at	gb:U49245.1 / DEE=Runan Gerangers / UG=Hs.78948 Rab gerangigerangitransferase, Deta Subunit / LESTERMINA / FEREMENA / FER
	gb:NM_002372.1 /DEF=Homo saptens memos and a same a same a same a relation of the same and same a same a same a class 2A, member 1 /DE XREF=g1:475-70-48:32965 mannosidase, alpha, class 2A, member 1 /DE XREF=G1-40-40-40-40-40-40-40-40-40-40-40-40-40-
at	gb:NM_006145.1 / DEF=Homo saplens near short and subfamily B, member 1 /FL=gb:BC002352.1 gb:NM_000125.1 gb:NM_0
	gb:BC000704.1 /DEF=Homo saptens, Lettaspars, Lettaspar
	gb.NM_018191.1 / DEF-Homo Saprens 1750-10-10-10-10-10-10-10-10-10-10-10-10-10
	gb:NM_014624.2 / DkF=Homo Saptems 35517 / UG=HS.275243 S100 calcium-binding protein Ab (calcyclin) / 12-35.200 / UG=HS.182490 leucihe-rich protein binding protein Ab (Jb_XREF=GI) / UG=HS.182490 leucihe-rich protein binding protein Ab (Jb_XREF=GI) / UB XREF=GI) / UB XREF
211971 s at	Consensus includes gp:Alousovo (Fig. 1200092) CONS. CDNA DKP2D564C046 (from clone DKP2D564C046). (FEA-mRNA /DB_XREF=g1:4500092) MRNA
	اتے :
1	protein FLJ22329 /FL=gb:NM 024656.1
	gb:NM_003983.1 /DEF=Homo sapiens solute carrier family / (cationic amino acidtransporter, y+ system), member 6 /DB_XREF=g1:450/U32 /UG=RB:11/012 /UG=RB:11/0
203580_s_at	(GENESAL AN) (INC. STATIONIC amino acid transporter, y+ system), member 0 (FLP9): DISTRIBLE IMAGE: 2228357 /UG-Hs. 75709 mannosh-6-phosphate carrier family 7 (cationic amino acid transporter, y+ system), MREF-est: tsl2d03.x1 /CLONE-IMAGE: 2228357 /UG-Hs. 75709 mannosh-6-phosphate
200900 s at	Consensus includes giralogolo (2015) (Free plane) (Free p
221652 s.at	gb:Ar274950.1 /DEF=Homo Sapieus rww. 2 mem., Protein PLJ13855.(FLJ13855), mRNA. /FEA=mRNA /GEN=FLJ13855 /PROD=hypothetical protein PrJ10637 /FL=gb:Ar274950.1
1 s 0 77777	8232
	signal cransducing anapton motif) 1 /DB_XREF=gi:4507248 /UG=Hs.15348/ signal transducing molecule (SH3 domainand TrAM motif) 1 /DB_XREF=gi:4507248 /UG=Hs.15348/
203544_s_at	cule (FACE-CESS) domain and ITAM motif) 1 /Firsts 1 /Firsts 1 /Firsts 1 /Firsts 1 /Firsts 2 /FEA-mRNA /PROD-Unknown (protein for MGC:5306) /DB_XREF=g1:12803030 /Cule (FACE-CESS) / DB_XREF=g1:12803030 /Cule (FACE-CESS) / DB_XREF=g1:128030 /Cule (FACE-CESS) / DB_X
221580_s_at	gp:BC0019/4:1 /DEF:NOORDED DEOLE N MCC5306 /FL=gb:BC001972.1 /VG=HS.301732 hypothetical protein MCC5306 /FL=gb:BC001972.1 /VG=HS.301732 hypothetical F70 /FFD=FST /DB XREF=FST:AV681579 /CLONE=GKBAFE05 /VG=HS.84084 amyloid beta precursor
202629_at	Consensus includes gainvoid. AFEA-mRNA /GEN-DDX18 protein 2 /FL-gb:AF017782.1 gb:NM 006380.1 Protein Cytoplasmic tail)-binding protein 2 /FL-gb:AF017782.1 gb:NM-DDX18 (Myc-regulated) (DDX18), mRNA. /FEA-mRNA /GEN-DDX18 protein Cytoplasmic tail)-binding protein 2 /FEA-mRNA /GEN-DDX18 (Myc-regulated) (ASP-Glu-Ala-ASPHIS) box polypeptide 18 (Myc-Rever 100555 DRADH (ASP-Glu-ASPHIS) box polypeptide 18 (Myc-Rever
205763 s at	Pron-DEADH (Asp-Glu-Ala-AspHis) box polypeptide 18 /DB_XKEF=91:12767557 Pron-DEADH (Asp-Glu-Ala-AspHis) box polypeptide 18 /DB_XKEF=est:tp93g08.x1 /CLONE=IMAGE:2206910 /UG=Hs.699 peptidylproly1 isom-regulated) /FL=gb:NML 006773.2
212337_at	Consensus includes 9D:Alb8/138 /FEA=E31 /DEA=E31 /DEA=E31 /DEA=E81 /DEA=E81 /DEA=E81 /FEA=ERNA FEA=ERNA FEA=E81 / FEA=E81 /DEA=E81 /DE
212244_at	1:4884111 /UG=H3.6283 DKFZF386F13 1299 /DB_XREF=est:7q83e05.x1 /CLC
222011_s_at	Consensus includes 90:5522000-1 Conzyme A thiolase) acetyltransferance and the AreFerit (PROD-NCK adaptor protein 1 / DB_XREFerit5453753 / Acetyltransferance and one NCK adaptor protein 1 (NCK1), mRNA. / PEA-mRNA / GEN-NCK1 / PROD-NCK adaptor protein 1 / DB_XREFerit55453753 /
.204725_s_at	gb:NX_U0b133.1 / DEF-LOAD Sup- UG-HS.54589 NCK adaptor protein 1 /FL-gb:NX_006153.1

1	The state of the s
200960 c at	gb:U09820.1 /DEF=Human helicase 11 (rADJ4n) make, compress (S. cerevisiae) homolog) /FL=gb:U09820.1 gb:NM 000489.1 gb:U/259: alloha thalassemiamental retardation syndrome X-linked (RAD54 (S. cerevisiae) homolog) /FL=gb:U08E:4185212 /UG=HS.37558 hypothetical prote-laloha thalassemiamental retardation syndrome X-linked (RAD54 (S. cerevisiae) homolog) /FL=gb:U08E:4185212 /UG=HS.37558 hypothetical prote-
20000	Consensus includes gb: PF940123 /FEA=EST /DB_XREF=g1: Lizocosos /LD_ALLE Consensus includes gb: PF940123 /FEA=EST /DB_XREF=g1: Lizocosos /LD_ALLE F=g1: Lizocoso /LD_ALLE F=g1: Li
203224_at	in Fight 149 (FER-gram Annatitis B virus X-associated protein 1 mRNA, complete cds. /FEA-mRNA /FROD=X-associated fire-in-
	2.521.00.WW. 001923.2 gb:L4032
208619_at	UG-HS.108327 damage-specific DNA binding protein 1 (12/KD) /FL-gb:urs.108327 damage-specific DNA binding protein 1 (12/KD) /FLA-CDS /GEN-HNRPR
	теорхосети
208765_s_at	/rws-BC001449.1 gb:AF000364.1 gb:NM_005826.1 RNA (BEA=mRNA /GEN=LOC51311 /PROD=T011-like receptor 8 (LOC51311), mRNA. /FEA=mRNA /GEN=LOC51311 /PROD=T011-like receptor 8
	gb:NM 016610.1 / DEF=Homo Saptens con 1 ch:NM 016610.1 db:AF245703.1 (/DB_XREF=gi:7706147 /
220832_at	UG-HS.272410 TOLL-like receptor # Fireguline receptor # FED-mann / GEN-TBPLI / PROD-TBP-like 1 / DB_XREF=91:4759233 / UG-HS. 2000 A.C. A.C. A.C. A.C. A.C. A.C. A.C. A
208198 s at	gb:NM 004865.1 /DEF=Homo Sapreus in increasing the control of the
1 0 00 00 00 00 00 00 00 00 00 00 00 00	nosphatidylinosicol .166982 phosphatidy
203011 Sac	th. BRD20712. I / DEF=Homo sapiens mRNA for KIAA0905 protein, compilete cus. / Lin-man.
210616 s at	VOG-HS. 70266 yeast Sec31p homolog /FL=gb:AB020712.1
211761_s_at	:2296820 /UG=Hs.324787 Solute Carrier
	Consensus includes guila Alaovia, inches 3 FELegh:NM 006933.1
213164_at	11/3 of Infortuce remarks by potherical protein FL311305 (F1311305), mRNA (FEMANICA (GEN-FLD11305) (F1311305)
219940_s_at	POSTAR TANGET 18922986 / UG-Hs 7049 hypothetical protein FLJ11305 / FL-GO:NM-INAGE:4476773 / UG-Hs.247324 hypothetical pro- FLJ11305 / DB_XREF=gi:8922986 / UG-Hs 7049 hypothetical protein FLJ11305 / UB-NAGE:4476773 / UG-Hs.247324 hypothetical pro-
	Consensus includes gb:B6254633 /FEATERS /FEATERS /CONSENSUS INCLUDES GB:B6254682 Apg12 (autophagy
203800_s_at	rein conserver in includes qb: BE965998 /FEA=EST / DB_XREF=gi:11770950 / DB_XREF=est:00103303281 / CLALL-
213026 at	12, S. cerevisiae)-like
	Consensus includes 9b:AL080099.1 /DEF=Humo Saptems AreF=g1:5262522 /UG=Hs.2057 uridine monophosphate synthetase (orocate phosphorizon)
215165 x at	Cara-Dar Sprower 12 (1988) Language and orotidine-5-decarboxylase) Lansferase and orotidine-5-decarboxylase)
. 45 0000	gb:NM_003113.1 /DEF=Homo sapiens nuclear antigen 5p100 /FL=gb:NG0618.1 gb:NM_003113.1 /NG=Hs.77617 nuclear antigen Sp100 /FL=gb:NG0618.1 gb:NM_003113.1
75 8 70000	gb:NM_001331.1 /DEF-Homo sapiens catenin (cadherin-associated protein), delta 1 /Frigos:NM_001331.1 delta 1 /Frigos:NM_001331.1 delta 1 /Frigos:NM_001331.1
	(cadherin-associated procenty, werea 1, 22, 22, 23, 23, 24, 25, 25, 25, 25, 25, 25, 25, 25, 25, 25
Ϋ.	gb: NM_002035.1"/DER=Homo sapiens follicular lymphoma variant translocation 1./PL=gb: NM_002035.1 1
202419_at	variant translocation 1. Described 1. Described describes 2 (HDAC2), mena. /FEA-mana /GEN-HDAC2 /FRUD-inscome ucaccety ass
201833 at	UG-HS.3352 histone deacetylase 2 /FL=gb:U31814.1 gb:NM 001527.1. //UG-HS.3352 histone deacetylase 2 /FL=gb:U31814.1 gb:NM 001527.1. //UG-HS.3352 histone deacetylase 2 /FL=gb:U31814.1 gb:NM 001527.1. //UG-HS.3352 histone deacetylase 2 /FL=gb:U31814.1 gb:NM 001527.1.
	Consensus includes government of the Est / Downer Street
217879_at	sion cycle 27 /FL=go:NM_Uutzoo.r

Tabelle 5: Gene aus Clusteranalyse 5

Affymetrix-	Affymetrix-Beschreibung der Seguenz in der Genebank Datenbank
interne	
Bezelchnung	
201466_s_at	gb:NM_002228.2 /DEF=Homo sapiens v-jun avian sarcoma virus 17 oncogene homolog (JUN), mRNA. /FEA=mRNA /GEN=JUN /PROD=v-jun avian sarcoma virus 17 oncogene homolog /DB_XREF=gi:7710122 /UG=Hs.78465 v-jun avian sarcoma virus 17 oncogene homolog /FL=cb:BC002646.1 cb:NM 002228.2
	gb:NM_000944.1 /DEF-Homo sapiens protein phosphatase 3 (formerly 2B), catalytic subunit, alpha isoform (calcineurin A alpha) PPP3GA), mRNA. /FEA-mRNA /GEN-PPP3GA /PROD-protein phosphatase 3 (formerly 2B), catalyticsubunit, alpha isoform (calcineurin A alpha) /DB_XREF=gi:6715567 /
202425_x_at	UG-Hs.272458 protein phosphatase 3 (formerly 2B), catalytic subunit, alpha isoform (calcineurin A alpha) /FL=gb:J05480.1 gb:L14778.1 gb:NM_000944.1 gb:AL353950.1
218566_s_at	gb:NM_012124.1 /DEF=Homo sapiens chord domain-containing protein 1 (CHPI), mRNA. /FEA=mRNA /GEN=CHPI /PROD=chord domain-containing protein 1 /PL=gb:AF192466.1 gb:NM_012124.1
214349_at	Consensus includes gb:AV764378 /FEA=EST /DB_XREF=g1:10922226 /DB_XREF=est:AV764378 /CLONE=MDSAGE03 /UG=Hs.163863 ESTs, Moderalely similar to POL2_HUMAN RETROVIRUS-RELATED POL POLYPROTEIN H.sapiens
204314_s_at	gb:NW_004379.1 /DEF-Homo saplens cAMP responsive element binding protein 1 (CREB1), mRNA. (FEA-mRNA /GEN-CREB1 /PROD-cAMP responsive element binding protein 1 /DB_XREF-g1:4758053 /UG-Hs.79194 cAMP responsive element binding protein 1 /FL=gb:NZ7691.1 gb:NM 004379.1
208753_s_at	gb:BC002387.1 /DEF-Homo sapiens, nucleosome assembly protein 1-11ke 1, clone MGC:8688, mRNA, complete cds. /FEA=mRNA /PROD=nhcleosome assembly protein 1-11ke 1 /DE_ab:BC002387.1 gb:AL1620681
	Consensus includes gb:AL031133 /DEF-Human DNA sequence from clone 281H8 on chromosome 6q25.1-25.3. Contains up to four novel genes, one with similarity to KIAA0323 and worm C30F12.1 and another with Ubiquitin-Like protein gene SWT3 (the latter in an introm of a novel gene). Con-
	ains up
215452_x_at	
204119_s_at	gb:U90339.1 /DEF=Human adenosine kinase short form nRNA, complete cds. /FEA=nRNA /PROD=adenosine kinase short form /DB_XREF=g4:1906010 / UG=HS-94382 adenosine kinase /FL=gb:U50196.1 gb:BC003568.1 gb:U90339.1 gb:UM_001123.1
201304_at	gb:NM_005000.2 /DEF=Homo sapiens NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 5 (13kD, B13) (NDUFA5), nuclear gene enchding mitochondrial protein, mRNA. /FEA-mRNA /GEN=NDUFA5 /PROD=NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 5 (13kD, B13) /FL-ab:BC000813.1 ab:NM 005000.2 ab:U51468.1 ab:U6-H5.0000.2
218131_s_at	gb:NM_017660.1 /DEF-Homo sapiens hypothetical protein FLJ20085 (FLJ20085), mRNA. /FEA-mRNA /GEN-FLJ20085 /PROD-hypothetical protein FLJ20085 /DG-Hs.118964 hypothetical protein FLJ20085 /FL-gb:NM_017660.1
201788_at	gb:NW_007372.1 /DEF=Homo sapiens RNA helicase-related protein (RNAHP), mRNA. /FERA=mRNA /GEN=RNAHP /PROD=RNA helicase-related protein / DB_XREF=gi:11321631 /UG=Hs.8765 RNA helicase-related protein /FL=gb:NM_007372.1 gb:AP083255.1
209780_at	gb:AL136883.1 /DEF=Homo sapiens mRNA; cDNA DKFZp434D166 (from clone DKFZp434D166); complete cds. /FEA=mRNA /GEN=DKFZp434D166 / PROD=bypothetical protein /DEF=g1:12053266 /UG=HS.128653 hypothetical protein DKFZp564F013 /FL=gb:AL136883.1
219375 at	gb:NM_006090.1 /DEF=Homo saplens cholineethanolaminephosphotransferase (CEPT1), mRNA. /FEA=mRNA /GEN=CEPT1 / PROD=cholineethanolaminephosphotransferase-/DB_XREF=qi:5174414 /UG=Hs.125031 cholineethanolaminephosphotransferase / FL=gb:AF068302.1 gb:NM_006090.1
209187_at	Consensus includes gb:AM516932 /FEA-EST /DB_XREF=gi:7154941 /DB_XREF=est:xq04a05.xl /CLONE=IMAGE:2748848 /UG-HS.16697 down-regulator of transcription 1, TBP-binding (negative cofactor 2) /FL-gb:BC002809.1
219279_at	gb:NM_017718.1 /DEF=Homo sapiens hypothetical protein FLJ20220 (FLJ20220), mRNA. /FEA-mRNA /GEN=FLJ20220 /FROD=hypothetical protein FLJ20220 /DE_KEF=gi:8923209 /UG=Hs.21126 hypothetical protein FLJ20220 /FL=gb:NM_017718:1
206976_s_at	H 144
215596_s_at	Consensus includes gb:AL163248 /DEF=Homo sapiens chromosome 21 segment HS21C048 /FEA=mRNA_2 /DB_XREF=gi:7717304 /UG=Hs.288773 zinc finger protein 294

	ned cus
216511_s_at	
201889_at	ء إس و
201435_s_at	Consensus includes grammars of the property of
212232_at	4 KI.
214329 x at	Consensus includes gb:AWW14434 (FEA-ES: 102-ALL: SECOND 1) AND TREFESSE: G108f09.x1 /CLONE=IMAGE:1855913 /UG=HS.204354 ras honolog gene similar to unknown H.sapiens
212099 at	Consensus includes gb:AI263309 /FEA=EST / DB_AKEF=91:30/212 / Consensus includes gb:AI263309 /FEA=EST / DB_AKEF=01 / Consensus framewore B /FEA=GRIN / GEN=KIAA0580 / PROD=KIAA0580 protein / Consensus B / FEA=GRIN / GEN=KIAA0580 / PROD=KIAA0580 protein /
213618 at	Consensus includes gb:AB011152.1 / DEF=Homo sapiens mana lot Arracoo Freezensus includes gb:AB011152.1 / DEF=Homo sapiens mana lot Arracoo Freezensus includes gb:AR0580 protein
216899 s.at	Consensus includes gb:AC003999 /DEF=Human PAC clone RP5-1139P1 from 7p15-p21 /FEA=CDS /DB_XREF=g1:2//2509 /UG-ns:2004 DEF=Human PAC clone RP5-1139P1 from factor 4E (RIF4E), mRNA. /FEA=mRNA /GEN=BIF4E /PROD=eukaryotic transla-
	gb:NM_001968.1 /DEF=Homo sapiens eukaryotic translation initiation factor 4E /FL=gb:MIS353.1 gb:NM_001968.1
201437_s_at	initiation factor 4E /DB_XREF=g1:45U3534 /UG=fa.155U0 cm.25.00 cm.
212122_at	
20866 s at	Consensus includes gb:BE866412 /FEA=ESI / DE_Arus -2. Consensus includes gb:BE866412 /FEA=ESI / DE_Arus - 1. Consensus includes gb:BE866412 /FEA=ESI / DECAMPS / GENECAPPS / FROD=Cytoskeleton associated protein cumorigenicity 13 (colon carcinoma) (HSp70-interacting protein) / Crasp) mRNA. / FEA=ERNA / GEN=CKAP2 / FROD=Cytoskeleton associated protein
	P-Homo sapiens cytoskeleton associated protesus a reservity and one of the protesus and other states of the protesus and other state
218252_at.	/DB_XREF=gi:8922641 /UG=Hs.24641 cytoskeleton associated protein 2 /Fir=gb:AL1364845.1 gb:Nn_Violate cds /FEA=CDS /DB_XREF=gi:8037944 /UG+Hs.283947 Homo
216384_x_at	gene, co
	Consensus includes gb:AK026451.1 / DEF = Homo Sapieus Common Consensus includes gb:AK026451.1 / DEF = Homo Sapieus Consensus gb:AK026451.1 / DEF = Homo Sapieus Consensus gb:AK026451.1 / DEF = Homo Sapieus Gb:AK026451.1 / DEF = Ho
20,000,000	is includes gb: Aa161486 /FEA=EST /DB XREF=gi:1735796 /DB XREF=est:2042005:54
204313_s_at	/FEA=EST
214257 s at	Consenses increment of the 1 inc
218595 s at	is hypothetical protein FLJ10359 (FLGDS) (18072.) (19072.
213128 g at	Consensus includes gb:AA527499 /FEA=EST /DB_XKEF=g1:4209300 /DL_ALALALALALALALALALALALALALALALALALALA
207719 x at	gb:NM_014812.1 /DEF=Homo sapiens KIAA0470 gene product (FL=gb:AB022657.1 gb:NM_014812.1 DB XREP=gi:7662141 /UG=Hs.25132 KIAA0470 gene product /FL=gb:AB022657.1 gb:NM_014812.1 DB XREP=gi:7662141 /UG=Hs.25132 KIAA0470 gene product /FL=gb:Admain_cich Admain_clone MGC:1125, mRNA, complete cds: /FEA=mRNA //
	3-Hs. 75104 RNA-binding protein S1, serine-rich domain
200060_s_at	
200634_at	M 005022.1
70 00000	gb:NM_004354.1 /DEF=Homo sapiens cyclin 62 (CCM62), huwn: /:
202//0_S_dL	

201008_s_at	Consensus includes gb:AA812232 /FEA=EST /DB_XREF=91:2881843 /DB_XREF=est:ob84h09.s1 /CLONE=IMAGE:1338113 /UG=Hs.179526 upregulated by 1.25-dihydroxyvitamin D-3 /FL=gb:NM_006472.1 gb:S73591.1
218534_s_at	
200993_at	Consensus includes gb:AA339270-/FEA-EST /DB_XREF=g1:3099183 /DB_XREF=est:oq31b02.s1 /CLONE=IMAGE:1587915 /UG=HS.5151 RAW binding protein 7 / FT=gb:AF098799.1 gb:NM 006391.1
214102_at	Consensus includes gb:AK023737.1 /DEF=Homo sapiens cDNA FLJ13675 fis, clone PLACE1011875, highly similar to Homo sapiens mRNA for KIAA0580 protein. /FEA=mRNA /DB_XREF=gi:10435758 /UG=Hs.287588 Homo sapiens cDNA PLJ13675 fis, clone PLACE1011875, highly similar to Homo sapiens mRNA for KIAA0580 protein
213704_at	19753 Lt
212367_at	Consensus includes gb:A1799061 /FEA-EST /DB_XREF=g1:5364533 /DB_XREF=est:we98a10.x1 /CLONE=IMAGE:2349114 /UG-HS.6048 FEM-1 (C elegans) homolog b /FL-gb:AF178632.1 gb:NM_015322.1 gb:AF204883.1
201873_s_at	gb:NW_002940.1 /DEF=Homo sapiens ATP-binding cassette, sub-family E (OABP), member 1 (ABCE1), mRNA. /FEA=mRNA /GEN=ABCE1 /PROD=ATF-binding cassette, sub-family E, member 1 /DE_XREF=gi:4506558 /UG=Hs.12013 ATP-binding cassette, sub-family E (OABP), member 1 /PI=-h-hNA nnooAn 1
218618_s_at	88
. 203689_s_at	/FEA=EST /DB_XREF=gi:5111325 /DB_XREF=est:w
201074_at	Consensus includes gb:AA593983 /FEA=EST /DB_XREF=gi:2409333 /DB_XREF=est:nn16d01.s1 /CLONE=INAGE:1084033 /UG=Hs.172280 SWISNF related, ma- trix associated, actin dependent regulator of chromatin, subfamily c, member 1 /FL=gb:U66615.1 gb:NM_003074.1
200778_s_at	Consensus includes gb:AI191427 /FEA+EST /DB_XREF=gi:3742636 /DB_XREF=est:qe48g03.x1 /CLONE=IMAGE:1742260 /UG-HS.155595 neural precursor cell expressed, developmentally down-regulated 5 /FL=gb:D28540.1 gb:NM_004404.1 gb:D63878.1
	gb:NM_000593.2 /DEF=Homo sapiens ATP-binding cassette, sub-family B (MDRTAP), member 2 (ABCB2), mRNA. /FRA=mRNA /GEN=ABCB2 /PROD=ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 2 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 3 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 3 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 3 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 3 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 3 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 3 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 3 /DB_XREF=gi:9665247 /UG=Hs.158164 ATP-binding cassette, sub-family B (MDRTAP), member 3 /DB_XREF=gi:966
213145_at	10. Date
218588_s_at	gb:NM_018691.1 /DEF=Homo sapiens chromosome 5 open reading frame 3. (C5ORF3), mNNh. /FEA=mNNh /GEN=C5ORF3 /PROD=hypothetical protein / DB_XREF=gi.8922068 /UG=Hs.166551 chromosome 5 open reading frame 3 /FL=gb:NM_018691.1
217834_s_at	gb:NM_006372.1 /DEF=Homo sapiens NSI-associated protein 1 (NSAPI), mRNA. /FFRA=mRNA /GEN=NSAPI /FROD=NSI-associated protein 1 / DB_XREF=gi:5453805 /UG=H8.155489 NSI-associated protein 1 /FL=gb:AR155568.1 gb:NM_006372.1
205038_at	14 21
203132_at	gb:NW_000321.1 /DBF=Homo sapiens retinoblastoma 1 (including osteosarcoma) (RBI), mRNA. /FEA-mRNA /GEN=RBI /PROD=retinoblastoma 1 (including osteosarcoma) /DB_XREF=g1:4506434 /UG=Hs.75770 retinoblastoma 1 (including osteosarcoma) /FL=gb:M33647.1 gb:M15400.1 gb:M28419.1 gb:N28419.1
211297_s_at	gb:L20320.1 /DEF-Human protein serinethreonine kinase stkl mRNA, complete cds. /FEA-mRNA /FROD-protein serinethreonine kinase / DB_XREF=gi:348242 /UG=Hs.184298 cyclin-dependent kinase 7 (homolog of Xenopus MOI5 cdk-activating kinase) /FL=cb:L20320.1
212375_at	/FEA=EST /DB_XREF=gi:129134
208986_at	Consensus includes gb:AL559478 /FEA=EST /DB_XREF=g1:12905019 /DB_XREF=est:AL559478 /CLONE=CSODJO13YM17 (5 prime) /UG=Hs.21704 transcription factor 12 (HTF4. helix-loop-helix transcription factors 4) /FL=gb:NB0627.1 gb:NB3233.1 gb:NM_003205.2
217956_s_at	gb:NM_021204.1 /DEF=Homo sapiens E-1 enzyme (MASA), mRNA. /FEA=mRNA /GEN=MASA /FROD=E-1 enzyme /DB_XREF=gi:10864016 /UG=HS.16442 E-1 enzyme //FL=gb:NM_021204.1 gb:AF13125.1
214870_x_at	Consensus includes gb:AC002045 /DEP=Human Chromosome 16 BAC clone CIT987SK-A-589H1 /FEA-mRNA_2 /DB_XREF=gi:2951945 /UG=Hs.251928 nuclear pore complex interacting protein.
209852_x_at	gb:BC001423.1 /DEF=Homo sapiens, Similar to proteaseome (prosome, macropain) 28 subunit, 3, clone MGC:1394, mRNA, complete cds. / FEA=mRNA /FROD=Similar to proteaseome (prosome, macropain) 28subunit, 3 /DB_XREF=gi:12655138 /UG=Rs.152978 proteasome (prosome, macropain)

rts.	activator subunit 3 (PA28 gamma; K1) /FL=gD:BC:001423;1 garcovert. 601859364F1 /CLONE=IMAGE:4069886 /UG=Hs.109643 polyadenylate
209063 x at	EA-EST /DB_XREF=gi:11164389 /DB_AAGE_=ESC: in 1 /PL=gb:AL136920.1
	ting
201177_s_at	gb:AF110957.1 gb:NM_005499.1 gb:AL130900.1
212264_s_at	Consensus includes gb:BE645850 /FEA=EST /DB XREF=91:39/0101 /DB_ANDELTON regulation 2, S. cerevisiae, homolog) 1 (SIRTI), mRNA. /
	וב שפרום
218878_s_at	-
202163 s at	
	1 /DEP=Homo sapiens KIAA0103 gene product (Arracios) 1 gb:NM_014673 61909 /UG-HS 154387 KIAA0103 gene product /FL=gb:D14659.1 gb:NM_014673
± .	Consensus includes gb:z14077.1 /DEF=H.sapiens mRNA for Yildr-El Piucell. / Linguistation factor /FL=gb:M7698.1 gb:M76541.1 gb:NM_003403.2
	034660 /DB_XREF=est:7d75b03.x1 /CLONE=IMAGE:3278/65 /UG-HS.108705 -J-7-2-1
	Consensus includes gb:T79584 /FEA=EST /DB_XREF=g1:598093 /DB_ANATATOLI gb:AF163473.1 gb:AF6254.1 gb:AF087438.1
202883_s_at	TOTMELY ZXV. TOTAL ZXV
202069_s_at	L=gb:NM_(
222303 at	Consensus includes gb:AV700891 (FEA=EST /DB_XREF=GI:10302862 /DB_XREF=EST:AV/00881 /CLONB-GAACASTO / REATS), mRNA, FEA=mRNA /
203102_s_at	GENEMBALZ / FACE-LIZE Deta-1, 2-N-acetylglucosaminyltransferase (rL=gD:Nn_cvc*vc; / CLONE=IMAGE:2237272 / UG=Hs.4014 KIAA0946 protein; (alpha-1,6-)-glycoprotein beta-1, 2-N-acetylglucosaminyltransferase; ts77a09.x1 / CLONE=IMAGE:2237272 / UG=Hs.4014 KIAA0946 protein;
212982 at	Consensus includes gb:AI611245 / FEA-E53 / VD-And S
200050_at	zinc
202430_s_at	gb.NM (021105.1 / DEF=HORD Septems Descripted scramblase 1 /FL=gb:NM 021105.1 gb:ABU00/40.1 gb:ABU005.1 pb:ABU00 ABU-1 BB:ABU00 ABU00 A
· 218757_s_at	3 212
214030_at	Consensus includes gorbeours, franche Company of the Company of th
218093_s_at	gb.NM_017664.1 /DEE=Homo Sapiems Proceedings
203486_s_at	Consensus includes gb: EL32373 FEL3 FEL3 FEL3 FEL3 FEL3 FEL3 FEL3 FEL
219303 at	gb:NM_024546.1 /DEF=Homo sapiens hypotherical protein FLJ13449 /FL=gb:AL136551.1 gb:NM_U44340.1 FLJ13449 /DB_XREF=g1:13375708 /UG=H9:10711 hypotherical protein FLJ13449 /FEA=mRNA /GEN=ZNF161 /PROD=zinc finger protein 161 /
202173 s at.	gb:NM_007146.1 /DEF=Homo sapiens zinc finger protein 161 /FL=gb:D28118.1 gb:NM_007146.1 DB_XREF=gi:6005967 /UG-Hs.6557 zinc finger protein
216903_s_at	Consensus includes gb: XX022897.1 / Ubr=non0 Septems / Ug=Hs.61628 calcium binding atopy-related automitigen / FEA-mRNA / DB_XREF=gi:10434244 / Ug=Hs.61628 calcium binding atopy-related automitigen CALC. / FEA-mRNA / DB_XREF=gi:10434244 / Ug=Hs.61628 calcium binding atopy-related automitigen calcium.

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205842_s_at	gb:AF001362.1 /DEF=Homo sapiens Jak2 kinase (JAK2) mRNA, complete cds. /FEA=mRNA /GEN=JAK2 /PROD=Jak2 kinase /DB_XREF=gi:3236321 / UG=Hs.115541 Janus kinase 2 (a protein tyrosine kinase) /FL=gb:NM_004972.2 gb:AF005216.1 db:AF001362.1 db:AF001362.1
212702_s_at	7 / DB_XREF=est:yz12f12.s1 /CLONE=IMAGE:282863 / mRNA, partial cds
201664_at	NA; cDNA DKF2p434F205 (from R=g1:6807670 /UG=Hs.50758 S
202060_at	gb:NM_014633.1 /DEF=Homo sapiens KIAA0155 gene product (KIAA0155), mRNA. /FEA=mRNA /GEN=KIAA0155 /PROD=KIAA0155 gene product / DE_KIREF=gi:7661949 /UG=Hs.173288 KIAA0155 gene product /FL=gb:NM_014633.1 gb:D63875.1
203177_x_at	ns transcription factor 6-like 1 (mitochond: uscription factor 6-like 1 (mitochondrialtr: or 6-like 1 (mitochondrial transcription fac
209748_at	S S
200660_at	202
222119_s_at	Consensus includes gb:AL117620.1 /DEF-Homo sapiens mRNA; cDNA DKFZp564K2364 (from clone DKFZp564K2364); partial cds. /FEA-mRNA / GEN-DKFZp564K2364 /PROD-hypothetical protein /DB_XREF-gi:5912200 /UG=Hs.284289 vitilion-associated protein VIR-1
208934_s_at	gb:AF342815.1 /DEF=Homo sapiens colorectal carcinoma-derived galectin-8 variant I mRNA, complete cds. /FEA-mRNA /PROD=colorectal carcinoma-derived galectin-8 variant1 /DB_XREF=gi:13249298 /UG=Hs.4082 lectin, galactoside-binding, soluble, 8 (galectin 8) / FI=gb:AF342815.1 gb:L78132.1 gb:AF074000.1 gb:NE_006499.1
211698_at	gb:AF349444.1 /DEF=Homo sapiens Rb- and p300-binding protein EID-1 mRNA, complete cds. /FEA=mRNA /PROD=Rb- and p300-binding protein EID-1 / DB_XNEF=g1:13549113 /FL=gb:AF349444.1
212053_at	Consensus includes gb:AK025504.1 / DEF-Homo sapiens cDNA: FLJ21851 fis, clone HEP01962. /FEA-mRNA / DB_XREF=gi:10438041 / UG-Hs.170218 KIAA0251 protein
209174_s_at	gb:BC000978.2 /DEF=Homo sapiens, hypothetical protein FLJ20259, clone MGC:5396, mRNA, complete cds. /FEA=mRNA /FROD-hypothetical protein FLJ20259 /PL=gb:BC000978.2 qb:NM 017730.1.
203476_at	gb:NM_006670.1 /DEF=Homo sapiens 574 oncofetal trophoblast glycoprotein (574), mRNA. /FEA-mRNA /GEN=574 /PROD=574 oncofetal trophoblast glycoprotein /DE_XREF=51:5729717 /UG=Hs.82128 574 oncofetal trophoblast glycoprotein /FE-cb:NM 006670.1
203007_x_at	gb:AF077198.1 /DEF=Homo sapiens lysophospholipase mRNR, complete cds. /FER=mRNR /PROD=lysophospholipase /DB_XREF=gi:4679009 /pG=Hs.12540 . lysophospholipase I /FL=gb:AF081281.1 gb:AF077198.1 gb:NR 006330.1 gb:AF291053.1
21,2721_at	Consensus includes gb:A1810380 /FEA=EST /DB_XREF=g1:5396946 /DB_XREF=est:wb87h10.x1 /CLONE=IMAGE:2312707 /UG=Hs.47274 Homo sapiens mRNA; cDNA DKFZp564B176 (from clone DKFZp564B176)
218461_at	.1 gb:\NY 016301.1
214218_s_at	Consensus includes gravessia, /rea=ESI /DB_XREF=g1:10301318 /DB_XREF=est:AV699347 /CLONE=GKCBPH07 /UG=Hs.83623.nuclear receptor.subfamily
1	gb:NM_020412.1 /DEF=Homo sapiens CHMP1.5 protein (CHMP1.5), mRNA, /FEA=mRNA /GEN=CHMP1.5 /PROD=CHMP1.5 protein /DB_XREF=gi:9966900 /UG=HS.42733
218178_s_at	
217851_s_at	gp:sw_ulov43.1 / DEFrando Saptens CGI-107 protein (LOC51012), mRNA. /FEA-mRNA /GEN=LOC51012 /FROD=CGI-107 protein /DB_XREF=gi:p705609 / UG=H3.3945 CGI-107 protein /FL=gp:AR151865.1 gb:NM_016045.1
202329_at	gb:NW_004383.1 /DEF=Homo sapiens c-src tyrosine kinase (CSK), mRNA. /FEA=mRNA /GEN=CSK /PROD=c-src tyrosine kinase /DB_XREF=g1:4758077 / UG=Hs:77793 c-src tyrosine kinase /FL=gb:NM_004383.1
203302_at	gb:NW_000788.1 /DEF=Homo sapiens deoxycytidine kinase (DCK), mRNA. /FEA=mRNA /GEN=DCK /FROD=deoxycytidine kinase /DB_XREF=g1:4503268 / UG=Hs.709 deoxycytidine kinase /FL=gb:M60527.1 gb:NM_000788.1
201949_x_at	Consensus includes gb:AL572341 /FEA=EST /DB_XREF=gi:12930514 /DB_XRRF=est:AL572341 /CLONE=CS0DI007xC13 (3 prime) /UG=Hs.76368 capping pro- Lein (actin filament) muscle Z-line, beta /FL=gb:NM_004930.1 gb:U03271.1

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	1988 - J. Amberdone hymothetical protein FLJ11210 (FLJ11210), mRNA. /FEA-mRNA /GEN-FLJ11210 /FROD-hypothetical protein
218096_at . I	gb:nM_UB351.1 / DEr-Bunn Supremarker hypothetical protein FLJ11210 /FL=gb:NM_UB351.1 PROD=vitiligo-associated protein VIT-1 PLJ11210 / DB_XREF=gi:8922941 / UG-HS:27842 hypothetical protein VIT-1 PROD=VIT1 / PROD=vitiligo-associated protein VIT-1
9 3256 25	gb:NM_018693.1 /DEF=Homo sapiens viiliigo-associated protein VIT-1 /FL=gb:NM_018693.1 gb:AF264714.1
203855_at	UG-EES. 2020 ALANDOS FACONOS OF SECRETOR REQUIRED FOR SPI transcriptional activation, subunit 3 (130kD) (CRSF3), MKWA. FEASTHOUSE SPIN. 0045FF FROM SPIN. 00
218846_at	for Spi transcriptional activation, subunit 3 (130kD) /FL=gb:AF135022.1 gb:AF10333.1 gb:AF-0333.1 gb:AF-0332.1 gb:AF-033.1 gb:AF-034.1 fb:AF-034.2 fb:
208988_at	
	n serine palmitoyitransierase (1662) muse, /UG=Hs.59403 serine palmitoyitransferase.
216202_s_ac	/DEF=Homo sapiens oligomycin sensitivity conferral protein oscp-like protein mona, partial cus- ferral proteinoscp-like protein /DB_XREF=gi:944964 /UG=Hs.76572 ATP synthase, H+ transporting, mi
216954_x_at .	in sensitivity conferring pro
212753_at	
	gb:NM_001776.1 /DEF=Homo sapiens ectonucieoside tribuospuene urpuospuenistatione (https://doi.org/101776.1
207691_x_at	triphosphate diphosphohydrolasel /DB_XREF=gi:4502666 /UG=Hs.205353 ectonucleoside triphosphohydrolasel /UG=Hs.20535 ectonucleoside triphosphoh
	apien)/DB
200967_at	gb:M63573.1 gb:NM_000942.1
201988 s at	Consensus includes garatecone. Priegb: AF039081.1 gb:NM 001310.1 ment binding protein-like 2 /Firegb: AF039081.1 gb:NM 001310.1
	gb: BC003090.1 / DEF-Homo sapiens, COP9 homolog, clone MGC:1297, mana, complete cds. / Fer-mana / Live Copy Con Complete Color / Fer-mana / Fer
202142_at	UG=HS. / 2133 COF9 Hountog first control of the con
218238_at	UG-HS.215766 GTP-binding protein /FL-gb:AF325353.1 gb:AF120334.1 gb:NM_UL2341.1 Cranscript variant B16, mNNA. /FEA-mNNA /GEN-OAS1 /
	gb.NM_002534.1 /DEF-Homo sapiens 2,5-oiigoadenylate synthetase 1 (**)-*
טחקקקט פ אד	/FL=gb:BC000562.1 gb:D00068.1 gb:NM 002534.1
2272-227007	/PEA-EST /DB_XREF-gi:11682003 /DB_XREF-est:naaU8DU5.X1 /CLONG-IMAGE:255577 / CC-255777 / CC-25577 / CC-255777 / CC-25577
202541_at	
221699_s_at	
	a di
202090_s_at	PL=gb:D55636.1 gb:NM_006830.1
202382_s_at	gp:nm_0034111 /Dir.com. Com. 100=Hs.278500 glucosamine-6-phosphate isomerase /FL=gb:NM_0054111 gp:ArU2351411 gb:ArU2351411 gb:ArU235141 gb:ArU2351
	/DEF=Homo sapiens transcription factor (p38 interacting protein, 1992) interacting protein) /FL= teracting protein) /DB_XREF=gi:8923734 /UG=Hs_171185 transcription factor (p38 interacting protein) /FL=
220408_x_at	gb:NM_017569.1 (EIF351), mRNA. /FEA=mRNA /GEN=EIF351
10000	gb:NM_003758.1 /DEF=Home Sapiens envaryor: transferent factor fac
208264_5_ac	sensus includes gb:AW024925 /FEA=EST /DB_XREF=gi:5878455 /DB_XREF=est:wu92g09.x1 /CLONE=IMAGE:232/332 /UG-ms:CU705
202872_at	ing,

α

pa306

	Consensus includes gb:AP131748.1 / DEF=Homo sapiens clone 25191 GTP-specific succiny1-CoA synthetase beta submit (SCS) man Sequence, Consensus includes gb:AP131748.1 / DEF=Homo sapiens clone 25191 GTP-specific succiny1-CoA synthetase betasubunit / DB_XREF=gi:4406563 / UG=Hs.247309 succinate-CoA ligase,
214835_s_at-	partial cds. //EA-mank fronts: or
205263_at	PROD=CARD-containing about to a system of the state of th
218512_at	gb:RM_018200.1 / DEF-Mome September 13291 hypothetical protein FLJ10881 /FL=gb:AF442340.1 gb:RM_01820.1 / DEF-Homo September 1 polyadenylate binding protein interacting protein 1, clone MGC:12360, mRNA, complete cds. / gb:BC005295.1 / DEF-Homo Sapiens, similar to polyadenylate binding protein 1 / DB_XREF=gi:13529010 / UG-Hs:109643 polyadenylate binding
210283_x_at	FEA=mRNA / PROD=SINILAR to polyacentrate
202724_s_at	gb:NM 002015.2 / DEF=Round Sapitans Corkhead box 01A (rhabdomyosarcoma) /FL=gb:AF032885.1 gb:UU2510.1 gb:NM_COLETT / UG=Hs.170133 forkhead box 01A (rhabdomyosarcoma) /FL=gb:AF032885.1 gb:UU2510.1 / UG=Hs.170133 forkhead box 01A (rhabdomyosarcoma) /FL=gb:AF032885.1 gb:UU2510.1 gb:NM_COLETT / UG=Hs.170133 forkhead box 01A (rhabdomyosarcoma) /FL=gb:AF032885.1 gb:UU2510.1 gb:UU25
	/FEA=EST /DB_XREF=gi:3735200 /DB_XREF=est:qd60D05.x1 /CLONE=IRM4E:1735041 /CC-H3.77495 UBX
212006_at	
222040_at	Consensus includes go:Alignov (Francisco Alignos) Consensus includes go:Alignos Consensus includes risonuclear
202373_s_at	complete
209301_at	M_000067.1
213264_at	Consensus includes go: NWZ-13-10 (12.00)
202863_at	gb:NM_003113.1 / DEF-man September 2010 / FLE-gb:NMO618.1 gb:NM_003113.1 UG=HS.71617 nuclear anticore anticore; / PRE-gb:NMO618.1 gb:NM_003113.1 / CLONE=IMAGE:2274675 / UG=HS.201605 ESTS, Moderately
217679_x_at	Consensus includes go: Alos Jose Alos Secure Contamination Warning ENTRY H. Sapiens similar to Alu8 HUMAN WALO SECURE CONTAMINATION WARNING ENTRY H. Sapiens Alberta to Homo sapiens mann contaminated to Homo sapiens conta
212802_s_at	Consensus includes gb: AkV23041.1 / Db: The Part of the Consensus includes gb: AkV2304.1 / Db: Transpa Dr. Consensus includes gb: AkV2312 (from clone Dr. Consensus of the Con
210346_s_at	S CLAS MANA
208848_at	gb:M30471.1 / DEF-Human class ii alcohol dehydrogenase 5 (class III), chi polypeptide /FL-gb:NM 000671.2 gb:M29872.1 gb:M30471.1 hase 3 / DB_XREF=gi:178133 /UG-Hs.78989 alcohol dehydrogenase 5 (class III), chi polypeptide /FL-gb:NM 000671.2 gb:M29872.1 gb:M30471.1 DB_XREF=gi:178133 /UG-Hs.110309 najor histocompati-
221978 at	
217526 at	Consensus includes gb:AI478300 /FEA-EST / DB_XKEF-91:4-71230 / DEA-EST / DB XKEF-91:4-71230 /
218322_s_at	gb:NM_016234.2 /DEF=HGmo Saptens long-chain rate acid coenzyme A ligase 5 /FL=gb:NM_01624.2 gb:nbc235525. coenzyme A ligase 5 /DE_RXREF=12 /UG=HS.11638 long-chain fatty acid coenzyme A ligase 5 /UG=HS.87497 butyrophilin,
212613_at	Consensus includes 95.4127122 (FEA-mRNA /GEN-FPMID / Subfamily 3, member A2 (FEA-mRNA /GEN-FPMID / MANNA /FEA-mRNA /GEN-FPMID / Subfamily 3, member A2
. 204566_at	gb:NM_003620.1 / DEF=Homo Sapieus process to magnesium-dependent, delta isoform / DB_XREF=gi:4505996 / UG=HS.100980 process pr

Tabelle 6: Gene aus Clusteranalyse 6

Affymetrix-	Beschreibung der Sequenz in der Genebank Datenbank
interne Be-	
zeichnung	
202018 s_at	gb:NM_002343.1 /DEF=Homo sapiens lactorransferin (Diff) (MASSO2.1 gb:NM_002343.1 /VG=HS.105938 lactotransferin /FL=gb:AF332168.1 gb:M350.1 gb:M3502.1 gb:NM_002343.1
	Consensus includes gb: AK000168.1 /DEF=Homo sapiens CDNA FLAZILIA 113. CONSENSUS FORMS Sapiens CDNA FLAZO161 fis, Homo sapiens CD24 signal transducer mRNA. /FEA=mRNA. /FEA=mRNA. / FIX- river files mRNA.
216379_x_at	DB XREF=qi:2810111 /DB_XREF=est:
. 209771 x at	Consensus includes guranoire, franciscome cluster 4 antigen) /FL=gb:X69397.1 //UG=Hs.286124 CD24 antigen (small cell lung carcinome cluster 4 antigen)
	gb:M33326.1 /DEF=Human nonspecific cross-reacting antigen (NVA) mank, complete cus. //PER=mRNA /GEN=NCA /PROD=non-specific cross reacting antigen (N B_XREF=91:189101)
206676_at	/UG=Hs.41 carcinoembryonic antigen-related cell adhesion molecule o /ru-guinalizati gen-HTm4 /PROD=IgE receptor beta
	gb:L35848.1 /DEF=Homo sapiens igE receptor Deta Chain (nims) mann, member 3 (hematopoietic cell-specific) / subnit /DB_XREF=gi:561638 /UG=HS.99960 membrane-spanning 4-domains, subfamily A, member 3 (hematopoietic cell-specific) /
210254_at	FL=gb:NM_006138.1 gb:L35848.1
203887 c at:	gb:NM_000361.1 /DEF=Homo sapiens thrombomodulin /FL=gb:NM_000361.1 UG-Hs. 2030 thrombomodulin /FL=gb:NIG-52.1 gb:NM_000361.1
35-0-1-0-1-0-1-0-1-0-1-0-1-0-1-0-1-0-1-0-	from clone RP11-209019 on intramestale is concern, cres (GW112 and KIAA4294) /FEA-mRNA /DB_XREF=gi:11182238
212768_s_at	differentially expressed in hematopoietic lineages
1	gb:NM 019058.1 /DEF=Homo sapiens nypothetical protein /FL=gb:AL136668.1 gb:NM_019058.1
20200	gb:NM_003564.1 /DEF=Homo sapiens transgelin 2 (IAGLNZ), mRNA. /FEA-mRNA /GEN=TAGLNZ /FRUD=Lransgelin 2 /DD_ANA.
200916_at	UG-HS. 75725 transgelin 2 /FLegb:BZ12611 do:Nex UG-BG from ranidly induced by IL-1 beta (FYX3), mRNR. /FEA-mRNR /GEN-FYX3 /
	gb:NM_002852.1 /DEF=Homo Saptens pencalnitered by IL-lbeta /DB_XREF=gi:4506332 /UG=Hs.2050 pentaxin-related gene, rapidly induced property induced by IL-lbeta /DB_XREF=gi:4506332 /UG=Hs.2050 pentaxin-related gene, rapidly induced by IL-lbeta /UG=Hs.2050 pentaxin-related
206157_at	by IL-1 beta /FL=gb:M311bb.1 gb:NN2-00252:1.
	gb:M63310.1 /DEF=Human 1,2-cyclic-incorror phosphodiesterase /DB_XREF=gi:178696 /UG=Hs.1378 annexin A3 /FL=gb:BC000871.1 PROD=1,2-cyclic-incorporate phosphodiesterase /DB_XREF=gi:178696 /UG=Hs.1378 annexin A3 /FL=gb:BC000871.1
209369_at	gb:m03320.1 gb:m2320.1 gs:m200.1 gs:m200.1 gs: 128826 f. DB_XREF=est:ts93d05.x1 /CLONE=IMAGE:2238825 /UG=HS:/594 soluce Consensus includes gb:AI631159 /FEA=EST /DB_XREF=q1:4682489 /DB_XREF=est:ts93d05.x1 /CLONE=IMAGE:2238825 /UG=HS:/594 soluce
202497_x_at	carrier family 2 (facilitated glucose transporter), memoer 3 / 22-92
	gb:NM_U01062.1 / DEF=ROMD Sapiens Critical Dinding protein, Rbinder family) /DB_XREF=g1:4507406 /UG=HS.2012 Franscobalamin I (vitamic Franscobalamin I (vitamic Franscobalamin I (vitamic Franscobalamin I) (vitamic Franscobalamin I)
205513_at	(vitamin B12 binding protein, k binder remary) (1789) (VITAMINA) (VITAMIN) PROD-haptoglobin (DB XREF-gi:4826761 /UG-Hs.75990 (VITAMINA) (DEN-HOMO Sapiens haptoglobin (HP), mRNA, (FEA-mRNA /GEN-HP /PROD-haptoglobin /DB XREF-gi:4826761 /UG-Hs.75990
206697_s_at	haptoglobin /FL=gb:K00422.1 gb:L29394.1 gb:NM_005143.1 haptoglobin /FL=gb:K00422.1 gb:L29394.1 gb:L293
202146_at	1
266 s at	L33930 /FEATURE / DEFINITION=HUMCD24B Homo sapiens CD24 signal transducer mRNA, complete cds and 3 region cross
100000	gb:MI8728.1 / DEF-Human nonspecific crossreacting antigen mRNA, complete cus. / Lin-Luck / Complete cu
21165/_ac	ובטרנינוט מוייים מייים ביי

205214_at [4] 205214_at [4] 91 207574_s_at gi	gb:NM_004256.1 /DEF=Homo sapiens serinethreonine kinase 17b.(apoptosis-inducing) (STK17B), mRNA. (FER=mRNA /GEN=STK17B / PROD=serinethreonine kinase 17b(apoptosis-inducing) /DB_XREF=gi:4758193 /UG=Hs:120996 serinethreonine kinase 17b
	(apoptosis-inducing) /Fb=gb:AB011421.1 gb:NM_004226.1
	gb:NM_015675.1 /DEF=Homo sapiens growth arrest and DNA-damage-inducible, beta (GADD45B), mRNA. /FEA-mRNA /GEN-GADD45B / PROD=DKFZFS66B133 protein /DB_XREF=gi:9945331 /UG=Hs.110571 growth arrest and DNA-damage-inducible, beta /FL=gb:AF090950.1 gb:NM_015675.1
206343_s_at //	gb:NW_013959.1 /DEF-Homo sapiens neuregulin 1 (NRG1), transcript variant SADF, mRNA. /FEA=mRNA /GEN=NRG1 /FROD=neuregulin 1 isoform SMDF //DB_XREF=gi:7669517 /UG=Hs.172816 neuregulin 1 /FL=cp;NM 013959.1 ch:141877 1
214146_s_at cc	Consensus includes gb:R64130 /FEA=EST /DB_XRRF=gi:836009 /DB_XREF=est:y118h03.s1 /CLONE=IMAGE.133637 /UG>Hs.2164 pro-platelet basic protein (includes platelet basic protein, beta-thromboglobulin, connective tissue-activating peptide III, neutrophil-activating peptide-2)
215716_s_at st	Consensus includes gb:L14561 /DEF=Homo sapiens plasma membrane calcium ATPase isoform 1 (ATP2B1) gene, alternative splice products, partial cds /FEA=mRNA_2 /DB_XREF=gi:4165324 /UG=Hs.78546 ATPase, Ca++ transporting, plasma membrane 1
9th 201179_s_at al	
201626_at /c	Consensus includes gb:BQ292233 /FEA=BST /DB XREF=gi:13050848 /DB XREF=est:602386668F1 /CLONE=INAGE:4515521 /UG=Hs.56205 insulin induced gene 1 /FL=cb:NM 005542 1
209728_at DI	gb:BC005312.1 /DEF-Homo sapiens, clone MGC:12387, mRNA, complete cds. /FEA-mRNA /PROD-Unknown (protein for MGC:12387)./ DB_XREF-gi:13529055 /UG-Hs.318720 Homo sapiens, clone MGC:12387, mRNA, complete cds /FL-cb:BC005312.1 ch:Mf642 1
212665_at UC	Consensus includes gb:AL556438 /FEA-EST /DB_XREF-gi:12899113 /DB_XREF-est:AL556438 /CLONE=CSODK004YI18 (3 prime) / UG-Hs.12813 DRF2P434J214 protein
92 GE GE GE GE GE GE GE	gb:NM_003118.1 /DEF=Homo sapiens secreted protein, acidic, cysteine-rich (osteonectin) (SPARC), mRNA. /FEA-mRNA / GEN=SFARC /PROD=secreted protein, acidic, cysteine-rich(osteonectin) /DB_XREF=gi:4507170 /UG=Hs.111779 secreted protein, acidic, cysteine-rich (osteonectin) /FL=gb:BC004974.1 gb:J03040.1 gb:NM_003118.1
201407_s_at ph	Consensus includes gb:AI186712 /FEA=EST /DB_XREF=gi:37351350,/DB_XREF=est:qe82£01.x1 /CLONE=INAGE:1745497 /UG=Hs.21537 protein phosphatase 1, catalytic subunit, beta isoform /FL=gb:NM_002709.1 gb:AF092905.1
gb //G 203505_at su	gb:AF285167.1 /DEF=Homo sapiens ATP-binding cassette transporter 1 (ABCA1) mRNA, complete cds. /FEA=mRNA /GEN-ABCA1 /PROD=ATP-binding cassette transporter 1 /DB_XREF=gi:9755158 /UG=Hs.211562 ATP-binding cassette, sub-family A (ABC1), member 1 /PL=gb:AF165281.1 gb:NR 005502.1 qb:AF285167.1
221802_s_at //U	Consensus includes gb:AU157109 /FEA-EST /DB_XREF-gi:11018630 /DB_XREF-est:AU157109 /CLONE-PLACE1006159 /UG-Hs.23740 KIAA1598 protein
201844_s_at /	Consensus includes gb:W84482 /FEA=EST /DB_XRRF=g1:1395613 /DB_XRRF=sst:zd89h07.s1 /CtCNR=IMAGE:356701 /UG=Hs.7910 RING1 and YY1 binding protein /FL=gb:AR179286.1 gb:AB029551.1 gb:NM_012234.1
206390_x_at /D	gp:NM_002619.1 /DEF=Homo sapiens platelet factor 4 (PF4), mRNR. /FEA=mRNR./GEN=PF4 /FROD=platelet factor 4 /PL=gb:MZ5897.1 gb:NM_002619.1
212569_at Co	Consensus includes gb:AV699744 /FEA=EST /DB_XREF=gi:10301715 /DB_XREF=est:AV699744 /CLONE=GKCEDF05 /UG=Hs.8118 XIAA0650 ordiein
211960_s_at //C	Consensus includes gb:BG261416 /PEA=EST /DB_XREF=g1:12711232 /DB_XREF=est:602373192F1 /UG=Hs.237955 hypothetical protein PR02706
gb // // // // // // //	gb:NM_022977.1 /DEF=Homo sapiens fatty-acid-Coenzyme A ligase, long-chain 4 (FACL4), transcript variant 2, mkNA. /FEA-mRNA /GEN-PACL4 /FROD=long-chain fatty-acid-Coenzyme A ligase 4,isoform 2 /DB_XREF=gi:12669908 /UG=HS.81452 fatty-acid-Coenzyme A ligase, long-chain 4 /FL=gb:NN_022977.1
221958_s_at //U	Consensus includes gb:AA775681 /FEA=EST /DB_XREF=g1:2835015 /DB_XREF=est:zf31a02.s1 /CLONE=IMAGE:378506 /UG=Hs.250746 hypothetical protein FLJ23091
204621_s_at . /U	Consensus includes gb:AI935096 /FEA=EST /DB_XREF=g1:5673966 /DB_XREF=est:wp13e10.x1 /CLONE=INAGE:2464746 /UG=HS-82120 nuclear receptor subfamily 4, group A, member 2 /FL=gb:NM_006186.1
gb / 201409_s_at //U	gb:NM_002709.1 /DEF=Homo sapiens protein phosphatase 1. catalytic subunit, beta isoform (PPPICB), mRNA. /FEA=mRNA /GEN=PPPICB /PROD=protein phosphatase 1. catalytic subunit, betaisoform /DB_XREF=gi:4506004 /UG=Hs.21537 protein phosphatase 1. catalytic subunit, beta isoform /FI=gb:NM_002709.1 gb:aF092905.1

201662_s_at	gb:D89053.1 /DEF=Homo sapiens mRNA for Acyl-CoA-synthetase 3, complete cds. /FEA=mRNA /PROD-Acyl-CoA synthetase 3 / CDB_XREF=gi:4165017 /UG=Hs.268012 fatty-acid-Coenzyme A ligase, long-chain 3 /FL=gb:NM 004457.2 db:D89053.1 db:AF116690.1
220987_s_at	gb:NM_030952.1 /DEF=Hamo sapiens hypothetical protein DKFZp434J037 (DKFZP434J037), mRNA. /FEA=mRNA /GEN=DKFZP434J037 /PROD=hypothetical protein DKFZp434J037 /DB_XREF=gi:13569921 /FL=gb:NM_030952.1
219607_s_at	gb:NW_024021.1 /DEF=Homo sapiens membrane-spanning 4-domains, subfamily A, member 4 (MS4A4), mRNA. /FRA-mRNA /GEN=MS4A4 /PROD=membrane-spanning 4-domains, subfamily A, member4 /DB_XREF=gi:13430865 /UG=HS.325960 membrane-spanning 4-domains, subfamily A, member 4 /FL=qb:AB013102.1 b:NM 024021.1 db:aP068288.1 db:NM 016650.1
201565_s_at	gb:NM_002166.1 (DEF=Homo saptens inhibitor of DNA binding 2, dominant negative helix-loop-helix protein (ID2), mRNA. /FEA-mRNA / GEN=ID2 /FROD=inhibitor of DNA binding 2, dominant negativehelix-loop-helix protein /DB_XREF=gi:4504501 of DNA binding 2, dominant negative helix-loop-helix protein /FI=qb:NM 002166.1 oh:DN protein /FI
213327_s_at	Consensus includes gb:AI820101 /FEA=EST /DB_XREF=G1:5439180 /DB_XREF=est:wg56d12.x1 /CLONE=IMAGE:2369111 /UG=Hs.42400 ubiquitin specific protease 12
203603_s_at	gb:NW_014795.1 /DEF=Homo sapiens zinc finger homeobox 1B (ZFHXIB), mRNA. /FEA=mRNA /GEN=ZFHXIB /PROD=zinc finger homeobox 1B / DB_XREF=sg1:7662183 /UG=Hs.34871 zinc finger homeobox 1B /PI=gb:NR 014795.1
217963_s_at	gb:NW_014380.1 /DEF=Homo sapiens p75NTR-associated cell death executor; ovarian granulosa cell protein (13kD) (DXS6984E), mRNA. /FEA=mRNA /GEN=DXS6984E /FROD=p75NTR-associated cell death executor; ovariangranulosa cell protein (13kD) /DB_XREF=g1:7657043 / UG=HS.17775 p75NTR-associated.cell death executor; ovarian granulosa cell protein (13kD) /FL=ch:NM (143R) 1 ch:ar18704 1
206710_s_at	gb:NM_012307.1 /DEF=Homo saplens differentially expressed in adenocarcinoma of the lung (KIAA0987), mRNA. /FEA=mRNA /GEN=KIAA0987 /PROD=differentially expressed in adenocarcinoma of the lung /DB_XRRF=g1:6912469 /UG=Hs.103839 erythrocyte membrane protein band 4.1-like 3 /FLFL=gb:AF069072.1 gb:NM_012307.1
204342_at	gb:NM_013186.1 /DEF=Homo sapiens hypothetical protein (DKFZp586G0123); mRNR. /FEA=mRNR /GEN=DKFZp586G0123 //PROD=hypothetical protein /DB_XREF=gi:9558726 /UG=Hs.24713 hypothetical protein /FL=qb;AL050209.1 gb;NM 013386.1
204393_s_at	gb:NM_001099.2 /DEF-Homo sapiens acid phosphatase, prostate (ACPP), mRNA. /FRA-mRNA /GEN-ACPP /PROD-prostatic acid phosphatase precursor /DB_XREF=g1:6382063 /UG=Hs.1852 acid phosphatase, prostate /FL=gb:MZ4902.1 gb:M34840.1 gb:NM_001099.2
200719_at	Consensus includes gb:BE964043 /FEA=EST /DB_XREF=g1:11767371 /DB_XREF=est:601657616R1 /CLONE=IMAGE:3875955 /UG=Hs:171626 transcription elongation factor B (SIII), polymeptide 1-like /FL=cb:NM 003197.2
202731_at	gb:NW_014456.1 /DEF=Homo sapiens programmed cell death 4 (PDCD4), mRNA. /FEA=mRNA /GEN=PDCD4 /FROD=programmed cell death 4 /FL=gb:U96628.1 gb:NM_014456.1
204094_s_at	iens KIAA0669 gene product 526 KIAA0669 gene product /
208690_s_at	gb:BC000915.1 /DEF=Homo sapiens, Similar to LIM protein; clone MGC:5344, mRNA, complete cds. /FEA=mRNA /PROD=Similar to LIM protein /DB_XREF=gi:12654194 /UG=Hs:75807 PDZ and LIM domain 1 (elfin) /FL=qb:BC000915.1
208979 at	gb:AF128458.1 /DEF-Homo sapiens nuclear receptor coactivator RAP250 mRNA, complete cds. /FEA-mRNA /PROD-nuclear receptor coactivator RAP250 /DB_XREF-g1:7140573 /UG-Hs.159613 thyroid hormone receptor binding protein /FL-gb:AF177388.1 gb:AF208227.1 gb:AF128458.1 gb:NM_014071.1 gb:AF171667.1 gb:AF245115.1
201772_at	gb:NM_015878.1 /DEF=Homo sapiens antizyme inhibitor (LOC51582), mRNR. /FEA=mRNR /GEN=LOC51582 /PROD=antizyme inhibitor / DB_XREF=gi:7706219 /UG=Ks.223014 antizyme inhibitor /FL=gb:D88674.1 gb:NM_015878.1
	gb:NM_003339.1 /DEF=Homo sapiens ubiquitin-conjugating enzyme E2D 2 (homologous to yeast UBC45) (UBE2D2), mRNA. /FEA=mRNA /GEN=UBE2D2 /PROD=ubiquitin-conjugating enzyme E2D 2 (homologousto yeast UBC45) /DB_XREF=g1:4507774 /UG=Hs.108332 ubiquitin-conjugating enzyme E2D 2 (homologous to yeast UBC45)
221060 s.at	Tru-garozzati. garam_Unasasi. garam_003266.1 DEF=Romo sapiez toll-like receptor 4 (TLR4), mRNR. /FEA=mRNR /GEN=TLR4 /PROD=toll-like receptor 4 /DB XRRF=d: 4507532 /MG=Hs 159239 foll-like receptor 4 /FEI-dh-MB9886 1 Jh-NR 003266 1
208878_s_at	gb:AF092132.1 /DEF=Homo sapiens PAKZ mRNA, complete cds. /FEA=mRNA /FROD=PAKZ /DB_XREF=gi:5138913 /VG=HS.284275 Homo sapiens PAKZ mRNA, 7complete cds /FL=gb:AF092132.1
212628_at	Consensus includes gb:BG292065 /FEA=EST./DB_XREF=gi:13050507 /DB_XREF=est:602386350F1 /CLONE=IMAGE:4515036 /UG=Hs.69171 protein kinase C-like 2
212928_at	Consensus includes gb:AI050331 /DEF-Human DNA sequence from clone 48613 on chromosome 6q22.1-22.3. Contains the part of a gene for a novel protein, the gene for KIAA0721 (NAP (Nucleosome Assembly Protein)

	3:- entering property, the TSPYL dene for TSPY-like (testis specific protei /FEA=mNNA_1
	اش
221568_s_at	highly similar to AF090900 Homo sapiens clone HULLS FROULS HANN FILTS (CLONE-IMAGE:342932)
201408_at	/UG=HS.21537 protein phosphatase 1, catalytic subunit, beta isoform /FL=gp:NM_U02/US.1 gp:Atoxxocx
	gb:NM_002053.1 /DEF=Homo sapiens guanylate binding protein 1, interferon-inducible, 67kD /DB_XREF=gi:4503938 //PEA=mRNA /GEN=GBP1 /PROD=guanylate binding protein 1, interferon-inducible, 67kD /DB_XREF=gi:4503938
202270_at	UG-HS.62661 guanylate binding protein 1, interferon-inducible, 0/AD / FEA-ERRNA /GEN-TIK1 /FROD-tousled-like kinase 1
202606_s_at	gb:NM_012290.1 / DEFENDANG Septems Coursied-like kinase 1 /FL=gb:AB004885.1 gb:NM_012290.1 gb:AFZ46215.1 / UG=HS.1892-1 / UG=HS.1892-1 / UB_XREF=g1:6912719 / UG=HS.1892-1 / UB XREF=g1:6912719 / UG=HS.1892-1 / UB XREF=g1:6910.1 / CLONE=IMAGE:895403
216037 x at	Consensus includes gb:AAbb4011 /FEA=E21 / DB_AADE_91:21 HMG-box Consensus includes gb:AAbb4011 /FEA=E21 (VG-HS:173638 transcription factor 7- like 2 (T-cell specific, HMG-box)
	gb:LZ1934.2 /DEF=Homo sapiens acyl-coenzyme A: cholesterol acyl.trans.texse.main.
221561_at	/UG-HS.14553 Steroi C-avyitimasspace of AMP-dependent, regulatory, type II, beta (PRKAK2B), mkWA. gb:Wm.002736.1 /DEF-Homo sapiens protein kinase, cAMP-dependent, regulatory, typeII, beta PRERF-914506064 // pen.mbwn /rFni=PRKAR2B /PROD=protein kinase, cAMP-dependent, regulatory, typeII, beta / nh.wk. nn7146.1
203680_at	/ug-Hs.77439 protein kinase, caMP-dependent, regulatory, type II, beta /ri-gb:hb.1120.1 go.m. or caMP-dependent, regulatory, type II, beta /ri-gb:hb.1745,
215111_s_at	/UG-Hs.114360 transforming growth factor beta-stimulated protein 150-22
'	gb:BC005122.1 /DEE=Homo saplems, ANT-12-2-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1
202211_at	protein 1 /FL=gb:BC005122.1 gb:AF111847.1 gb:NM_014570.1
221918_at .	Consensus includes gb:AI742210 /FEA=EST / DB XREF=eg1:5110498 / DB XREF=est:W712064 / CLONE=DCAAUDO5 / UG=HS.9456 SWISNF related, matrix
•	Consensus includes gb:AV712064 /FEA=EST / DB_ARDE =g1:L012130 / L2
213251_at	ident regulator of
209186_at	Example 1 Jose Louis Des Line Slow twitch 2 /FL-gb:W23114.1 Example 14-3-3 protein /DB_XREF=gi:4096984 /UG=Hs.79474 Example 14-3-3 protein JDE XREF=gi:4096984 /UG=Hs.79474 Cample 14-3-3 protein mRNA, complete cds. /FEA=mRNA /PROD=14-3-3 protein JDE XREF=gi:4096984 /UG=Hs.79474
210996_s_at	
	Consensus includes gb:AA8/3600 /FEA-651 /DD_CONSENSUS includes gb:AA8/3600 /FEA-651 /FEA-6
213624_at ·	like phosphodiesterase
1	gb.NM_00315.1 / DEF=nomb Saprems Screens Screen Screens Screens Screens Screens Screen
2000 - COMO	Consensus includes gb:BG492602 /FEA=EST /DB_XREF=gi:13454114 /DB_XREF=est:001220505011 /Consensus includes gb:BG492602 /FEA=EST /DB_XREF=gi:13454114 /DB_XREF=gi:17705613 /
217215_s_ac	gb.NW 016048.1 /DEF=Homo saptens CGI-111 protein (LOC51015), mRNA. /FEA-mRNA /GEN-LOC31013 /FAND-CGI /FEA-GRNA /FEA-MRNA /GEN-LOC31013 /FAND-CGI /FEA-GRNA /
218170_ac	
210293_s_at	siae) homolog B /DB XREF=gi:13477148 /UG=Hs.173497 Sec23 (S. cerevisiae) homolog B /FL=gb:BCU534.1 homolog B /DB XREF=gi:13477148 /UG=Hs.173497 Sec23 (S. cerevisiae) homolog B /FL=gb:BCU534.1 homolog B /DB XREF=gi:13477148 /UG=Hs.173497 Sec23 (S. cerevisiae) homolog B /FL=gb:BCU534.1
221493 at	gb:AL136629.1 /DEF=Homo Saprens mark) Company Topy-like /FL=gb:AL136629.1 prop=hypothetical protein /DB_XREF=gi:12052783 /UG=Hs.278479 TSPY-like /FL=gb:AL136629.1

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	Consensus includes go:bE905050 / FEM-ES1 / DE_ANDE_S1:2000 / PEN-ES1 / DE ANDE ANDE ANDE ANDE ANDE ANDE ANDE A
213300_84	.2684 /FEA=EST /DB_XREF=gi:9771329 /DB_XREF=est:601067180F1 /CLONE=IMAGE:3433371 /06=EB:3320 curriform
201016 at	tion initiation factor 1A /FL=gb:BC000793.1 gb:L18960.1 gb:NM_001412.1
	jation-associated (mm), muni. /ibn-man. 07 /UG-HS.79889 monocyte to macrophage differentiation-associated
203414_at	FL=gb:NM_012329.1
213510_x_at	TOL TL132 /GEN-FLJ20725 /PROD-hypothetical protein
	gb:NM_017943.1 /DEF=Homo sapiens hypothetical protein FL460/23 (suscense); makes fractions fract
218539_at	DB_XREF=g1:8923650 /UG=HS.15467 hypothetical protein FL/20/25 /FL=gD:NM_UL/343.1. NRNA. /PEA=mRNA /GEN=FR02577 /PR02577 (PR02577), mRNA. /FEA=mRNA /GEN=FR02577 /PR0D=hypothetical protein PR02577
,	
218172_s_at	(DKFZP5640123), 1
202538 s_at	gb:NM_014043.1 / DEF=Homo Saptens DAFEFSTANIZ Protein /FL=gb:AF151842.1 gb:AL080122.1 gb:NM_014043.1 DB_XREF=gi:7661633 / UG=H8.11449 DKFZP5640123 protein /FL=gb:AF151842.1 gb:AL080122.1 gb:NM_014043.1
	abunt Sfu alpha isolothi mana) compress
221504 s.ac	subunit SFD alpha isoform /DB_XREF=g1:6563195 /UG=Hs.19575 CGI-11 protein /FL=gb:AF132945.1 gb:AF132945.1 gb:AF112204.1 gb:AF11204.1 gb:AF
	139.1 /DEF-Homo sapiens mRNA for KLAAUS6/ proceim, partial
212213 x at	DB_XREF=gi:3043657 /UG=HS.147946 optic atrophy 1 (autosomal dominant) / CEA=mRNA /DB XREF=gi:6807841 /
	ייין מייין אייין
216652_s_at	EF-Human DNA sequence from
	cellular
:	retinatoenyde-Dinding process; do see see see see see see see see see se
221472_at	OB XREF=g1:9650676 /UG=Hs.272168 tumor differentially expressed i /fu-gr:05.000 5 sumilar to Homo sapiens i
	Consensus includes gb:AK002207.1 / DEF noun Saprens Core. 12010.1 / KIAA0610
212526_at	protein. /FRA=mRNA /DB_XREF=gi:7023938 /UG=Hs.118087 XIAAUb1U protein. /FRA=mAGE:1084659 /UG=Hs.78825 matrin 3 /
200624 s at	Consensus includes gb:AA5/1095 /FEA=EST / DB_AAGE-91:233301 / JEB-05:00 / JEB 1 gb:AB018266.1
	Consensus includes gb:BE748755 /FEA=EST /DB_XREF=g1:10162747 /DB_XREF=est:bUL5/193311 /CLUME-IMAGE:300013 /
201091 s at	11ke protein 1 /FL=gb:AF136630.1 gb:NM_016587.1
	18378 /FEA=EST /DB_XREF=gi:4085584 /DB_AKEF=gsc:gozumou.xl /cucal_mass_mass_mass_mass_mass_mass_mass_ma
217863 at	AspHis) box binding protein 1 /FL=gb:AF077951.1 gb:AF167160.1 gb:NM_016166.1
	sensus includes gb:AI5
202804 at	te, sub-family C (CFTRNRP), member 1 /FL=gb:L05628.1 gb:NM 004996.2
	Consensus includes gb:AU150824 /FEA=EST /DB_XREF=g1:L1012343 /Lb_Arkf=cst.noilcol. /
214198_s_at	
	gb:NM_016141.1 /DEP=Homo sapiens dynein light chain-A (LOCSIL43), maya. / LACSIL43.1
217976_s_at	(/DB_XKEF=91:170303Z /

	UG=Hs.266483 dynein light chain-A /FL=gb:AF078849.1 gb:NM_016141.1	
	gb:AF306508.1 /DEF-Homo sapiens SUMO-1 specific protease PKSG6 mRNA, complete cds. /FEA-mRNA /PROD=SUMO-1 specific protease PKSG6 /	. ,
202318 s at	DB_XREF=gi:11096243 /UG=Hs.2/119/ SUMO-1-Specific protects /firsts mana, complete cds. /FEA=mRNA /GEN=SFRS3 /PROD=pre-mRNA splicing factor (SFRS3) mana, complete cds. /FEA=mRNA /GEN=SFRS3 /PROD=pre-mRNA splicing factor	factor
208673_s_at	/ DB_XREF=gi:5531903 /UG=HE.167460 splicing factor, arginineserine-rich 3 /FL=gb:BC000914.1 gb:AF107405.1 DB_XREF=gi:5531903 /UG=HE.167460 splicing factor, arginineserine-rich 3 /FL=gb:BC000914.1 gb:AF107405.1	uţ
	gb:NM_022771.1 / DEF=Homo sapiens hypothetical protein FLuizuos (FLuizuos), muxm. / Linearia (School of Particula)	•
218268_at	DB_XREF=gi:12232450 /UG-HS:48827 hypothetical protein functions from Protein is: linid desaturase), clone McC:5079, mRNA, complete cds.	cds. /
	gb:BC000961.2 /DEF-Homo sapiens, degenerative spermatocyte (nominos receptivase) /DB_KREF=91:12803018 /UG-Hs.185973 degenerative spermato-FEA-mRNA /FROD-degenerative spermatocyte (homolog Drosophila;lipid desaturase) /DB_KREF=91:12803018 /UG-Hs.185973 degenerative spermatocyte	jermato-
209250_at	cyte (homolog Drosophila; lipid desaturase) /FL=gb:BC000961.2	23
	GD:NM_015957.1 /DEF=Homo sapiens CGI-29 protein (LOC51074), mRNA. /FEA=mRNA /GEN=LOC31074 /FROJ-CGI-22 protein	
218698 at	/UG=Hs.104058 CGI-29 protein /FL=gb:AF132963.1 gb:NM_015957.1	

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Patentansprüche

- Array bestehend aus Oligo- oder Polynukleotidsonden,
 die immobilisiert auf einem festen Träger aufge bracht sind, dadurch gekennzeichnet, dass auf der
 Oberfläche Sequenzen einer Auswahl oder aller der
 in den Tabellen 1-6 genannten selektiven Monozyten Makrophagen-Gene gebunden sind.
- 2. Array nach Anspruch 1, dadurch gekennzeichnet, dass gegebenenfalls zusätzlich weitere Gene verwendet werden, von denen bekannt ist, dass sie in jeder Zelle exprimiert werden und zur Grundausstattung einer Zelle gehören.
 - 3. Array nach Anspruch 1 und 2, dadurch gekennzeichnet, dass mit den genannten Genen komplementäre RNA auf der Oberfläche des Arrays gebunden ist zum inversen Nachweis über die in den Tabellen 1-6 dargestellten Gene oder Gensequenzen.
 - 4. Array nach Anspruch 1 bis 3, dadurch gekennzeichnet, dass die Gene, deren Teil- und Oligomersequenzen krankheits- und nebenwirkungsrelevante selektionierte Gene der rheumatoiden Arthritis oder anderer chronisch entzündlichen Erkrankungen vor und nach anti-TNF-Therapie sind.
- 5. Array nach Anspruch 1 bis 4, dadurch gekennzeichnet,
 dass die Gene, deren Teilsequenzen und Oligomersequenzen krankheitsspezifisch regulierte Gene des
 Monozyten/Makrophagen-Zellsystems sind.

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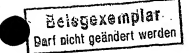
- 6. Array nach Anspruch 1 bis 5, dadurch gekennzeichnet, dass auf der Oberfläche gegebenenfalls auch Allele, Derivate und/oder Splicingvarianten der Gen- bzw. Genteilsequenzen und Oligomersequenzen vorliegen.
- 7. Array nach Anspruch 1 bis 6, dadurch gekennzeichnet, dass es auf der Oberfläche Gensequenzen enthält, die mindestens eine Teil-Sequenzidentität von 80 % in den Protein-kodierenden Abschnitten der mRNA besitzen.
- 8. Array nach Anspruch 1 bis 7, dadurch gekennzeichnet, dass die Oberfläche der Träger mit reaktiven Gruppen, Metallverbindungen oder Legierungen beschichtet ist.
- 9. Array nach Anspruch 1 bis 8, dadurch gekennzeichnet, dass die Gene oder Gensequenzen durch Spottingverfahren von cDNA, Immobilisierungs-verfahren und Syntheseverfahren von Oligomeren oder spiegelbildlich in Form von RNA aufgebracht sind.
- 10. Verwendung des Arrays nach Anspruch 1 bis 9 mit Proben, die zum Nachweis Fluoreszenzfarbstoff-, Enzym-, Protein- oder radioaktiv markiert sind und eine Verstärkung zulassen.
- 11. Verwendung des Arrays nach Anspruch 1 bis 9 mit Proben, dadurch gekennzeichnet, dass die Verstärkung der Signale über gekoppelte alkalische Phosphatase, Peroxidase, Biotin Digoxigenin-, Proteinmoleküle, (Edel-)Metallchelate oder Beads erfolgt.

- 12. Verwendung des Arrays nach Anspruch 1 bis 9 mit Proben, dadurch gekennzeichnet, dass zur zusätzlichen Verstärkung der Signale Streptavidin, (Edel-)Metallchelate, Beads oder Antikörper eingesetzt werden.
- 13. Verwendung des Arrays nach Anspruch 1 bis 9 zum inversen Nachweis festphasengebundener Total-RNA oder messenger-RNA.
- 10 14. Verwendung des Arrays nach Anspruch 1 bis 9 zur Messung der Monozyten/Makrophagen-Aktivierung oder der Entzündungsaktivität im Blut oder im Zellgewebe.

- 15. Verwendung des Arrays nach Anspruch 1 bis 9 zur

 Feindiagnostik sowie zur Früherkennung von entzündlichen Erkrankungen und der rheumatoiden Arthritis.
- 16. Verwendung des Arrays nach Anspruch 1 bis 9 zur Verfolgung von Nebenwirkungen bei der anti-TNF-Therapie von entzündlichen Erkrankungen und der rheumatoiden Arthritis.
 - 17. Verwendung des Arrays nach Anspruch 1 bis 9 zur Überwachung der Therapie und Erstellung einer Prognose bei entzündlichen Erkrankungen und der rheumatoiden Arthritis.
 - 18. Verwendung der Arrays nach Anspruch 1 bis 9 zur Identifizierung von pharmazeutischen Targets bei entzündlichen Erkrankungen und der rheumatoiden Arthritis.

- 19. Verwendung der Gene oder Gensequenzen nach Tabelle 1-6 zu Einzelgennachweisverfahren, vorzugsweise reverse Transkriptions-PCR (RT-PCR).
- 5 20. Verwendung der Gene oder Gensequenzen nach Tabelle 1-6, dadurch gekennzeichnet, dass sie mit einer Markierung oder einer Reporterfunktion ausgestattet sind.
- 10 21. Verwendung der Gene oder Gensequenzen nach Tabelle 1-6 zum reversen Nachweis festphasengebundener Total-RNA oder messenger-RNA in einem RNA-Array mit bis zu 500 Gewebs- und/oder Blutproben.



Zusammenfassung

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Die Erfndung betrifft einen Array bestehend aus Oligodie immobilisiert auf einen oder Polynukleotidsonden, festen Träger aufgebracht sind. Das Array ist dadurch charakterisiert, dass auf der Oberfläche Sequenzen einer Auswahl oder aller der in den Tabellen 1-6 genannten selektiven Monozyten-Makrophagen-Gene gebunden sind. Array ermöglicht die Diagnose der rheumatoiden Arthritis und anderer chronisch entzündlicher Erkrankungen, begleitende Analyse der Behandlungseffektivität und die der bei Nebenwirkungen von Überwachung Tumornekrosefaktor (TNF)-Therapie und somit die Auswahl der für den jeweiligen Patienten mit rheumatoider Arthritis am wirkungsvollsten Therapie. Die vorliegende Erfindung betrifft ferner einen Nukleinsäure-Array zur Prognose und zur Entwicklung neuer anti-TNF gerichteter Pharmaka oder solcher Pharmaka, die in dessen Regelkreis eingreifen.

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